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- 6 Cloned human serum albumin gene.
- Disclosed are a synthetic human serum albumin gene, plasmids containing the gene, and microorganisms transformed by those plasmids.

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CLONED HUMAN SERUM ALBUMIN GENE

This invention relates to a method for synthesising a human serum albumin gene. This invention further relates to a plasmid containing a cloned human serum albumin gene and a microorganism transformed with such a plasmid.

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Human serum albumin (sometimes referred to

10 hereinafter as HSA) is the major protein component of
plasma. The protein is produced in the liver and is
primarily responsible for maintaining normal osmolarity
in the bloodstream. It also is capable of binding and
transporting various small molecules via the blood.

HSA is administered in various clinical situations. Shock and burn victims, for instance, usually require doses of HSA to restore blood volume and thus ameliorate some of the symptoms associated with trauma. Persons suffering from hypoproteinemia or erythroblastosis fetalis also are likely to require treatment with serum albumin.

To date, HSA is produced primarily as a by-product from the fractionation of donated blood. A drawback to this is that the cost and supply of blood can vary widely. The blood also may contain undesirable agents such as hepatitis virus. It therefore would be advantageous to develop an alternative source of HSA.

It accordingly is an object of this invention to produce human serum albumin in microorganisms. It is a further object of this invention to so produce HSA economically. It also is an object of this invention to

develop a cloning procedure that can be applied to other serum proteins.

Brief Description of the Figures

Figure 1 shows a partial restriction map of a fulllength HSA cDNA clone isolated by the procedures described herein.

Figure 2 shows the DNA sequence of the 5'+3' strand of the non-coding and coding regions of the full length HSA cDNA, as well as the amino acid sequence specified by the DNA sequence.

Figure 3 shows an A_{260} profile of sucrose gradient fractions of mRNA. Fraction group B was used as the template in the synthesis of HSA cDNA.

Figure 4 shows pGX401, a recombinant plasmid containing a full length HSA cDNA insert.

Figure 5 shows the DNA sequence in the region of codon 97 for HSA sequences derived from three different human livers.

According to one aspect of the present invention, we provide a synthetic human serum albumin gene. The term "synthetic" as used herein should be understood to include DNA sequences produced by use of recombriant DNA techniques and/or chemical synthesis.

In accordance with the present invention, a novel human serum albumin (HSA) gene has been cloned and bacterial expression of the gene is described. The nucleotide sequence of the full length HSA gene and the amino acid sequence of the polypeptide specified by that gene also are reported herein.

The procedure more fully described hereinafter which has been used to prepare an HSA-producing microorganism can be divided into the following stages: (1) obtaining HSA mRNA from a suitable source, e.g. by recovery and isolation of the HSA mRNA from HSA producing cells, (2) in vitro synthesis of complementary DNA (cDNA), using the mRNA as a template and conversion of the cDNA to the double-stranded form and (3) insertion of the double-stranded

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cDNA into a suitable cloning vector and transformation of microbial cells with that cloning vector. The procedures described herein resulted in the preparation of a "full-length" cloned HSA cDNA.

Eukaryotic genes are contained in the chromosomal DNA of cell nuclei. This chromosomal DNA exists in a compact nucleoprotein complex called chromatin.

Bukaryotic chromosomal DNA contains intervening sequences (introns) within the coding sequences (exons), which would not permit correct expression in bacteria. For this reason a preferred method for producing contiguous coding blocks of a particular protein involves the use of messenger RNA (mRNA). Messenger RNA has a ribonucleotide sequence corresponding to the gene of interest without the introns and conveniently can be recovered from eukaryotic cells that produce the protein specified by the gene.

Human serum albumin mRNA can be recovered in useful quantities from human liver cells. The HSA mRNA produced by the liver cells is complementary to one of the two strands of the HSA gene and may be employed as a template for the synthesis of complementary DNA (cDNA) as hereinafter described. To effectively utilize the mRNA for the synthesis of cDNA, it advantageously is recovered from the cells in relatively pure form. The guanidine thiocyanate/guanidine hydrochloride extraction procedure described by McCandliss et al., Methods in Enzymology 79:51 (1981), advantageously may be used to recover and purify the HSA mRNA. RNA is inherently less stable than DNA, and is particularly subject to degradation by ribonucleases that are present in the cells. Therefore, mRNA recovery procedures generally employ means for rapidly inactivating any ribonucleases which are present.

In general, recovery of total RNA is initiated by disrupting the cells in the presence of a ribonuclease-inactivating substance. Disruption of the cells may be

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accomplished by subjecting the cells to a lysing reagent, freezing/thawing, or mechanical disruption; preferably a combination thereof. A mixture of guanidine thiocyanate and a reducing agent, such as mercaptoethanol, has been found to function effectively as a ribonuclease inactivator (McCandliss, et al., supra).

After disruption of the cells, the solid cell debris is removed, e.g. by centrifugation, and the RNA is precipitated from the resulting clarified solution.

10 Precipitation is effected by known techniques, such as adding a water-miscible alcohol, e.g. ethanol, to the solution in a precipitating amount. The RNA then is resuspended in a guanidine hydrochloride solution and precipitated with ethanol for two successive cycles. At this point the RNA is undegraded and free of proteins and DNA.

The next step is the separation of mRNA from the total precipitated RNA. Human serum albumin mRNA is polyadenylated, therefore, it readily can be separated from non-adenylated RNA by affinity chromatography with oligodeoxythymidylate (oligo dT) cellulose (Aviv, H., et al., Proc. Natl. Acad. Sci. USA 69: 1408 (1972); McCandliss, et al., supra). Total RNA can be applied to a column in an approximately 0.5 M NaCl containing solution. Under these conditions only poly A+ RNA binds to the oligo dT cellulose and can be removed specifically by washing the column in a salt free solution.

To enrich the preparation for HSA mRNA, the poly A+RNA can be fractionated according to size by sucrose gradient centrifugation. Activity of the RNA in the various gradient fractions can be verified by in vitro translation in a reticulocyte lysate (Pelham, H., et al. Eur. J. Biochem. 67:247 (1976)) and by electrophoretic analysis of the protein products (Laemmli, U., Nature 227:680 (1970)).

Once a poly A+RNA fraction able to synthesize proteins the size of HSA has been isolated, it can be used to provide a template for cDNA synthesis. This procedure involves enzymatically constructing double-stranded DNA, which has a nucleotide base pair sequence identical to the coding sequence of the original chromosomal gene. The cDNA does not contain any noninformational segments (introns) within the coding region which might be present in the eukaryotic gene, and thus can ultimately be transcribed and translated in prokaryotic systems.

Synthesis of HSA cDNA employs the enzymes reverse transcriptase, Klenow fragment of DNA polymerase I and S1 nuclease (Racian, D., et al., Proc. Nat. Acad. Sci. USA 15 73:2191 (1976); McCandliss, R., et al., Methods in Enzymology 79, p. 601 (1981)). Reverse transcriptase catalyzes the synthesis of a single strand of DNA from deoxynucleoside triphosphates on the mRNA template. The poly r(A) tail of the mRNA permits oligo (dT) (of about 20 12 to 18 nucleotides) to be used as a primer for cDNA synthesis. The use of a radioactively-labelled deoxynucleoside triphosphate facilitates monitoring of the synthesis reaction. Generally, a @32p-containing deoxynucleoside triphosphate advantageously may be used 25 for this purpose. The cDNA synthesis generally is conducted by combining the mRNA, the deoxynucleoside triphosphates, the oligo (dT) and the reverse transcriptase in a buffered solution. This solution is incubated at an elevated temperature, e.g., about 40-50°C, for a time sufficient to allow formation of the cDNA copy, e.g. 30 about 5-20 minutes. The conditions of the reaction are essentially as described by Kacian, D.L., et al., supra. After incubation, disodium ethylenediaminetetraacetic acid (hereinafter EDTA) is added to the solution, and the 35 solution is extracted with phenol:chloroform (1:1 by

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vol.). The aqueous phase is advantageously purified by gel filtration chromatography, and the cDNA-mRNA complex in the eluate is precipitated with alcohol.

The mRNA can be selectively hydrolyzed in the . presence of the cDNA with dilute sodium hydroxide (about 0.1 M) at an elevated temperature, e.g., about 60-80°C for about 15-30 minutes. Neutralization of the alkaline solution and alcohol precipitation yields a single-stranded cDNA copy.

The single-stranded cDNA copy has been shown to have 10 a 5'-poly (dT) tail, and to have a 3' terminal hairpin structure, which provides a short segment of duplex DNA (Bfstratiadis, A., et al., Cell, 7, 279 (1976)). hairpin structure can act as a primer for the synthesis of a complementary DNA strand. Synthesis of this 15 complementary strand is conducted using the Klenow fragment of DNA polymerase I (Klenow, H., et al., Eur. J. Biochem., 22, 371 (1971)) in a reaction mixture containing the deoxynucleoside triphosphates. The duplex cDNA recovered by this procedure has a 3' loop, resulting 20 from the 3' hairpin structure of the single-stranded cDNA copy. This 3' loop can be cleaved by digestion with the enzyme, S1 nuclease, using essentially the procedure of McCandliss et al., Methods in Enzymology 79:601 (1981). The S1 nuclease digest may be extracted with phenol-25 chloroform, and the resulting cDNA precipitated from the aqueous phase with alcohol.

The intact double-stranded DNA (about 2000 base pairs) corresponding to a human serum albumin gene can be isolated by, for example, sucrose gradient centrifugation, using the procedure of McCandliss supra p. 51. In order to determine the sizes of the DNA in the sucrose gradient, aliquots of the gradient fractions are electrophoresed in a polyacrylamide gel with molecular weight markers. The resulting gel is first stained with

ethidium bromide to visualize the markers and then autoradiographed to detect the radioactive cDNA. The fractions of the gradient containing DNA molecules larger than 1000 base pairs are pooled and the DNA is precipitated with ethanol.

5 For purposes of amplification and selection, the double-stranded cDNA gene prepared as described above is generally inserted into a suitable cloning vector, which is used for transforming appropriate host cells. Suitable cloning vectors include various plasmids and 10 phages, with plasmids being preferred in this case. The criteria for selecting a cloning vector include its size, its capability for replicating in the host cells, the presence of selectable genes, and the presence of a site for insertion of the gene. With respect to its size, the 15 vector is advantageously relatively small, to permit large gene insertions, and so as not to divert large amounts of cellular nutrients and energy to the production of unwanted macromolecules. The vector also includes an intact replicon which remains functional 20 after insertion of the gene. This replicon preferably directs the desired mode of replication of the plasmid, i.e., multiple copies or a single copy per cell, or a controllable number of copies per cell. Genes specifying one or more phenotypic properties, preferably antibiotic 25 resistance, facilitate selection of transformants. The insertion site is advantageously a unique restriction site for a restriction endonuclease. A cloning vector meeting all of these criteria is the plasmid pBR322. The cDNA can be conveniently inserted into this plasmid by a 30 homopolymeric tailing technique. Homopolymer tails are added to the 3'-hydroxyl groups of the human serum albumin double-stranded cDNA gene, by reaction with an appropriate deoxynucleoside triphosphate, in the presence of terminal deoxynucleotidyl transferase. The plasmid is 35

opened by digestion with the appropriate endonuclease, and complementary homopolymer tails are added to the 3'hydroxyl groups of the opened plasmid, using the homopolymeric tailing technique. Appropriate reaction conditions have been described for the addition of dC 5 residues to ds cDNA (McCandliss, R., et al., page 601 supra; Roychoudhury, R., et al., Nucleic Acids Research 3:101 (1976)) and of dG residues to PstI treated pBR322 (Maeda, S., Methods in Enzymology 79:607 (1981)). preferred embodiment, however, the molar excess of dXTPs 10 to 3' ends is in the range of 3000 to 5000. Progress of the reactions is monitored until the chain length is approximately 15. The tailed cDNA and plasmids are recovered, e.g., by phenol extraction followed by alcohol precipitation. The homopolymeric ends of the two DNAs 15 are complementary and will anneal together under appropriate conditions to yield a recombinant plasmid containing the HSA gene (Maeda, S., Methods in Enzymology 79:611 (1981)).

A suitable strain of <u>E.coli</u> may be transformed with this recombinant plasmid, using essentially the method of Lederberg, <u>J. Bacteriology</u> 119:1072 (1974) and be maintained indefinitely.

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Cenerally, several hundred to several thousand

clones are produced by these procedures and can be
screened for the presence of the HSA gene with, for
example, rat serum albumin cDNA. A nick translated
(Maniatis, T., et al., Proc. Natl. Acad. Sci. USA 72:3961
(1975)) rat cDNA having 85% homology with human cDNA can

be used to hybridize to plasmid cDNA attached to
nitrocellulose filters (Grunstein, M., et al., Proc.
Natl. Acad. Sci. USA 72:396 (1975), Southern, E.M. J.
Mol. Biol., 98:503 (1975)). In this procedure, DNA from
each colony (or from groups of colonies) is fixed to
discrete zones of a nitrocellulose filter and denatured.

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Alternatively, the DNA can be electrophoresed in a gel prior to fixing on a filter. A solution of the radioactively labeled rat cDNA is applied thereto under hybridizing conditions. Unhybridized rat cDNA is washed from the filter, and colonies containing DNA to which the rat cDNA hybridized are identified by autoradiography. One positive clone was identified but found to be an incomplete HSA cDNA by DNA sequencing. A portion of this HSA cDNA was then nick translated in order to rescreen the entire bank of clones. Ninety positive hybridization signals were thus obtained.

Positive clones may be cultivated on suitable growth media to obtain ample quantities of cells from which to extract the plasmid DNA. The plasmid DNA is extracted, using conventional techniques, such as disruption of the cells, followed by phenol extraction, and alcohol precipitation. The plasmid and chromosomal DNAs may be separated, e.g. by electrophoresis or cesium chloride equilibrium centrifugation. Plasmid DNA containing inserts of about 1500 to 2000 base pairs are selected for further characterization.

The cloned gene can be excised from the plasmid DNA and then characterized by sequencing analysis (Sanger, F., et al., Proc. Natl. Acad. Sci USA 74:5463 (1977); Maxam, A., et al., Proc. Natl. Acad. Sci. USA 74:560 (1977)).

By these procedures a prepro-HSA clone has been isolated. An E. coli HB101 culture transformed with the plasmid containing this prepro-HSA gene has been deposited with the U.S. Department of Agriculture Northern Regional Research Laboratory in Peoria, Illinois, as NRRL No. B-15784. A diagnostic partial restriction map of this HSA gene insert is shown in Figure 1 of the drawings and Figure 2 shows the 5'-->3'

strand of the non-coding and coding regions, along with the amino acid sequence specified by the gene.

The cloned prepro-HSA coding sequence consists of 2050 base pairs excluding the oligo dC tails added to the cDNA. The gene has noncoding regions at the 5' end (base pairs 1-31) and at the 3' end (base pairs 1858-2050). The 5' end of the coding region (32-103 base pairs) includes a 24 amino-acid leader (an 18- amino-acid "pre" sequence followed by a 6-amino-acid "pro" sequence) and the mature human serum albumin protein is specified by the region from base pair number 104 to base pair number 1858.

As used in Figure 2 and elsewhere herein, the abbreviations have the following standard meaning:

	A	. =	deoxyadenyl
15	T	=	thymidyl
	G	=	deoxyguanyl
	С	=	deoxycytosy1
	GL	z =	glycine
	AL	4 =	alanine
20	VAI	; =	valine
	LE	u =	leucine
	IL	<u> </u>	isoleucine
	SE	R =	serine
	THI	R =	threonine
2 5	Pfi	E =	phenylalanine
	TY	R =	tyrosine
	TR	P =	tryptophan
	CY	S =	cysteine
	ME	T =	methionine
30	AS	P =	aspartic acid
	GL	0 =	glutamic acid
	LY	s =	lysine
	AR	.G =	arginine
	HI	s =	histidine
35	PR	0 =	proline

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GLN = glutamine

ASN = asparagine

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It will be appreciated that because of the degeneracy of the genetic code, the nucleotide sequence of the gene can vary substantially. For example, portions or all of the gene could be chemically synthesized to yield DNA having a different nucleotide sequence than that shown in Figure 2, yet the amino acid sequence would be preserved, provided that the proper codon-amino acid assignments were observed. Baving established the nucleotide sequence of the human serum albumin gene and the amino acid sequence of the protein, the gene of the present invention is not limited to a particular nucleotide sequence, but includes all variations thereof as permitted by the genetic code.

It is believed that the amino acid sequence set forth in Figure 2 and claimed herein represents a genomic HSA allele that is widespread in the human population, in contrast to the sequences previously published in the scientific literature. Polymorphism is known for HSA. Protein electrophoresis has revealed over twenty genetic variants of HSA (Weitkamp et al., Ann. Hum. Genet. London 36:381 (1973)). Two differing amino acid sequences have been reported previously. See Lawn, R.M., et al., Nucl. Acids Res. 9:6103 (1981) and Dugiaczyk, A., et al., PNAS 79:71 (1982). The DNA sequence of Figure 2 differs from each of these published sequences. Although some of the differences occur in third base position of codons or in the noncoding regions, and as such do not cause amino acid changes, conflicting nucleotide sequence data suggest different amino acids at positions 97 and 396. In Figure 2, the amino acid represented by codon 97 (GAG) is glutamic acid. The same was reported by Lawn, et al., supra. Dugiaczyk, however, reported that codon to be GGG (glycine). Codon 396 in

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Figure 2, also is designated GAG (gluatmic acid).

Dugiaczyk reported the same; however, Lawn reported codon

396 to be AAG (lysine). Thus, each of the three DNA

sequences would encode a different polypeptide. Example

IV below sets forth the procedures followed to determine
that these differences represented true protein

polymorphism and not merely experimental artificats.

The present invention has been described in connection with the use of <u>B. coli</u> as the bacterial host for recombinant DNA containing the HSA gene, but skilled molecular biologists will appreciate that other gramnegative bacteria, such as <u>Pseudomonas</u>; gram-positive bacteria, such as <u>Bacillus</u>; higher unicellular organisms, such as yeasts and fungi, and mammalian cells can be employed for cloning and/or expression of the BSA gene.

The invention is further illustrated by reference to the following examples, which are not intended to be limiting.

EXAMPLE I

Isolation of HSA mRNA from Human Liver Tissue

Messenger RNA (mRNA) was isolated from human liver tissue taken from a 10-year-old accident victim. Extreme care was taken throughout the procedures to avoid ribonuclease contamination of the mRNA preparation. These measures included the use of new, sterile laboratory glassware, treatment of solutions with diethylpyrocarbonate when appropriate, followed by autoclaving, keeping the preparation cold when possible and using gloves to avoid contact of the preparation with skin.

Frozen human liver tissue (10.5 grams) was homogenized in 210 mls lysis solution (4M guanidine thiocyanate/0.1M Tris-HCl, pH 7.5/0.1M 2-mercaptoethanol) using a Virtis homogenizer. Cellular debris was pelleted by

centrifugation at 8750 rpm, 4°C, for 10 minutes in a Sorvall GSA rotor, and the supernatant was transferred to a new centrifuge bottle. To the supernatant were added 0.04 volume 1M acetic acid and 0.5 volume 95% ethanol. After 2 hours at -20°C, the mixture was centrifuged at 5 7500 rpm, 10 minutes, 4°C and the pellet resuspended in 50 mls wash solution (6M guanidine hydrochloride/10mM Na₂·EDTA, pH 7.0/10mM dithiothreitol.) Centrifugation at 5500 rpm, 10 minutes, pelleted particulate debris, and the supernatant was transferred to a new centrifuge 10 bottle. To the supernatant were added 0.04 volume 1M acetic acid and 0.5 volume 95% ethanol. After 2 hours at -20°C, the mixture was centrifuged at 7200 rpm 20 minutes. The pellet was resuspended in 20 mls wash solution, and 0.04 volume 1M acetic acid and 0.5 volume 15 95% ethanol were added. The mixture was kept at -20°C for 12 hours, then centrifuged at 8,000 rpm for 10 minutes at 4°C in a Sorvall SS-34 rotor. The pellet was resuspended in 15 mls sterile distilled H20 (dH20) and extracted with an equal volume of (4:1) chloroform: 20 butanol. The aqueous phase was transferred to a fresh tube and 0.1 volume 2.4 M sodium acetate and 2.5 volumes 95% ethanol were added. After 2.5 hours at -20°C, the RNA was pelleted by centrifugation and the pellet was resuspended in 2 mls sterile dH_0 0). A total of 19.2 mg 25 RNA was recovered.

mRNA was then separated from the total RNA using generally, the oligo(dT)-cellulose affinity chromatography procedure described in Aviv et al.. supra and McCandliss, et al., supra. A column of 5 grams oligo(dT)-cellulose was washed with one column volume 0.1M NaOH to denature any ribonuclease present, then equilibrated with high salt buffer (10mM Tris-HCl, pH 7.4/0.5M NaCl/0.5% sodium dodecyl sulfate). The total RNA preparation, dissolved in two mls dH₂O above, was

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heated at 70°C for 1 minute, then cooled on ice to room temperature. Next, 0.1 volume 5M NaCl, 0.04 ml 0.5M Tris-HCl, pH 7.5, and 0.1 ml 10% sodium dodecyl sulfate (SDS) were added to the RNA. 8 mls high salt buffer were then added to the RNA and the solution was applied to the 5 column with a flow rate of about 10 drops/minute. After the sample had passed through, unbound RNA was washed from the column with high salt buffer. Practions (1/2 ml each) were collected and the optical density at 260 nm (A260) of each fraction was measured in a 10 spectrophotometer. The column was washed until the ${\tt A_{260}}$ readings of fractions dropped below 0.05. Undesired RNA was further washed from the column with low salt buffer (10mM Tris-HCl, pH 7.4/0.2M NaCl/0.1% SDS) and fractions 15 were collected as above until the A260 had dropped to 0.05.

Next, the mRNA was eluted from the column with elution buffer (10mm Tris-HCl, pH 7.4/1mm EDTA/0.1% SDS) and 1 ml fractions were collected until the A_{260} was less than 0.05. The first 15 fractions (those having the highest OD_{260} readings) were pooled and the mRNA was precipitated by adding 0.1 volume 2.4M sodium acetate and 2.5 volumes 95% ethanol, and placing at -20°C for 12 hours. The eluted mRNA was then pelleted by centrifugation and resuspended in 800 µl elution buffer. After heating the resuspended pellet at 70°C for 90 seconds then cooling on ice; 0.1 volume 5M NaCl and 0.05 volume 10% SDS were added.

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The eluted mRNA prepared above was then further

purified by passage over a second oligo(dT)-celluose
column. A column containing 0.1 gram oligo(dT) cellulose
was washed with NaOH, then with high salt buffer as
previously described. The RNA was applied to the column
and fractions were collected with high salt, low salt,

and elution buffers as with the first column. The peak

fractions from the elution buffer step were pooled and the twice-purified mRNA was precipitated and pelleted as before.

The mRNA was then size-fractionated on a 12-ml sucrose gradient as described in McCandliss et al.,

Methods in Enzymology, 79, pp. 56-58. A 5-20% sucrose gradient was prepared in gradient buffer (0.02M sodium acetate, pH 5.6) and chilled at 4°C for 3 hours. 100µg of the mRNA was resuspended in 100µl gradient buffer, heated at 80°C for 2 minutes, quick-cooled in an ice bath, then layered on top of the gradient. A second 5-20% gradient had E. coli 16 and 23S rRNA (100µg total) loaded on it to serve as molecular weight markers.

The two gradients were centrifuged in a Beckman SW40 rotor at 38,000 rpm for 12.5 hr at 4° C. Fractions of about 0.5 ml were then collected and the A_{260} measured (fraction $\frac{1}{2}$ 1 is that collected from the bottom of the gradient tube.) The A_{260} peak was divided into 6 groups of fractions, groups A through F as shown in Figure 3. The fractions in each group were pooled and the mRNA precipitated with 0.1 volume 2.4 M sodium acetate and 2.5 volumes 95% ethanol.

Praction groups containing mRNA which encodes protein of the size expected for HSA were identified by in vitro translation using a rabbit reticulocyte lysate kit (available from Bethesda Research Laboratories and used according to manufacturer's instructions) supplemented with ³⁵S methionine. A reaction mixture for each fraction group contained the components necessary for translation of the mRNA into radioactively-labeled proteins which were visualized by electrophoresis on a 12.5% polyacrylamide/SDS gel, followed by fluorography.

The fluorogram showed a prominent protein band of the size expected for HSA (68,000 daltons) among the translation products of fraction groups B and C. Group B

had a much lower percentage of protein products in undesirable low molecular weight range so the mRNA in group B was chosen for use as a template in the synthesis of cDNA.

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EXAMPLE II Synthesis of HSA cDNA

Generally, the cDNA synthesis procedure of McCandliss et al., Methods in Enzymology, 79, pp. 601-607 (1981) was used. Incorporation of a radioactively labeled deoxynucleotide allowed monitoring of the synthesis and calculation of yields at each step.

The first strand of cDNA was synthesized on the mRNA template, using oligo-dT as a primer, as follows.

Prepared mix and kept on ice:

	Frebarca man and mob-	
15	0.5 M Tris-HC1, pH 8.3	20µ1
	1.4 M KCL	10µ1
	0.25M MgCl ₂	8 µ l
	0.05M datp, pH 7.0	_ 2µ 1
	0.05M TTP, pH 7.0	2 µ 1
20	0.05M dCTP, pH 7.0	2µ1
20	0.05M dGTP, pH 7.0	2 u 1
	0.01M dithiothreitol	4µ 1
	sterile distilled 820	45 µ l
	aqueous label, $\alpha^{32}P-dCTP$ (10 μ Ci/ μ 1)	5µ1
	aqueous label, d Fadel (1000),	100µ1
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Added remaining components:

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	oligo(dT) $_{12-18}$ (250 μ g/ml)	20 µ l
	actinomycin D (500 µg/ml, aqueous)	16µl
	10µg mRNA, "B" fraction	20 µ l
5	sterile dH ₂ O	37µ1
	*AMV reverse transcriptase (16u/µl)	7 µ 1
	Total volume:	200µ1

*Avian myeloblastosis virus (AMV) reverse transcriptase is kept at -80°C and thawed briefly to add 10 as last component

The reaction mixture was kept on ice 5 minutes and 2 pl were removed and counted in ASC scintillation fluid in order to determine the specific activity of the dCTP. The reaction mixture was then incubated 10 minutes at 46°C. 20 pl 0.2M EDTA pH 8.0 was added to stop the reaction, and the mixture was then extracted with an equal volume (1:1) phenol:chloroform.

0.14 volume 80% glycerol was added and sample was chromatographed on a 0.7 x 17 cm. Sephadex G-100 column. Once the sample had entered the column, G100 buffer (10mM Tris-HCl, pH 8.0/1mM EDTA/100mM NaCl) was added to the column and 5-drop (about 275µ1) fractions were collected. The radioactive fractions were "Cerenkov counted" and the cDNA fractions comprising the peak counts per minute were pooled. The mRNA/cDNA hybrids were precipitated by adding 0.1 volume 2.4M sodium acetate and 2.5 volumes 95% ethanol, placing in a dry ice/ethanol bath for 30 minutes, then pelleting by centrifugation at 10,000 rpm, 4°C, for 20 minutes. The pellet was resuspended in 300µl 0.1M NaOH and heated at 70°C for 20 minutes to hydrolyze the RNA, leaving single-stranded cDNA. 30µ1 1M HCl were added to neutralize the solution. The DNA was precipitated by adding 5µg tRNA, 1/10 volume 2.4M sodium acetate, and 2.5 volumes 95% ethanol, placing in a dry

ice-ethanol bath 10 minutes, and centrifuging in a microfuge 10 minutes at 4°C.

The pellet was resuspended in the following mix:

40µl 0.5M potassium phosphate, pH 7.4

8µ1 0.25M MgCl₂

2µl 0.1M dithiothreitol

1µ1 0.05M dATP, pH 7.0

1µ1 0.05M dCTP, pH 7.0

1µ1 0.05M dGTP, pH 7.0

1µ1 0.05M TTP, pH 7.0

124µl sterile dH20

178µ1

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Next, added 22µl DNA polymerase I Klenow fragment (5µ/µl, available from Boehringer-Mannheim.)

The reaction mixture was then incubated in a 15°C water bath for 12 hours. 20µl 0.2M EDTA pH 8.0 was added to stop the reaction and the mixture was extracted with an equal volume (1:1) phenol:chloroform. 0.14 volume glycerol was added to the aqueous phase.

The sample, which now contains double-stranded cDNA, was run over a Sephadex G100 column and the peak cDNA fractions were pooled and precipitated as before. The double-stranded DNA has a 3' "hairpin loop" as previously described, which was removed with S1 nuclease as follows. The pellet was resuspended in 72 µl sterile distilled water and then 18 µl 5X S1 buffer (1M NaCl/0.25M sodium acetate, pH 4.5/5mM ZnSO₄/2.5% glycerol) were added. An enzyme mix was prepared by adding 2.5 µl (50 units) of S1 nuclease (20µg/µl) to 47.5 µl 1X S1 buffer. 10µl of enzyme mix was added to the 90µl DNA solution then incubated at 37°C 20 minutes. Addition of 20 µl 0.2M sodium BDTA stopped the reaction, and the reaction mixture was extracted with an equal volume (1:1) phenol:chloroform. The aqueous phase was

loaded onto a 5-25% sucrose gradient and spun at 38,000 rpm 17.5 hours 5°C in an ultracentrifuge.

One-ml fractions were collected and "Cerenkov counted." Fractions were pooled with fractions 1-6. 7-9. and 10-12 comprising the 3 pools. Fraction #1 was the 5 fraction 'taken from the bottom of the gradient. DNA was precipitated by adding 0.1 volume 2.4M sodium acetate, 1-2 µg tRNA, and 2.5 volumes 95% ethanol to each pool, then placing them at -20°C overnight. The DNA was pelleted by centrifugation at 25K for 30 minutes at 4°C. After 10 slightly dessicating pellets, the DNA from each pool was resuspended in 200 pl dB20 and precipitated again with ethanol and sodium acetate. Pellets were resuspended in 22v1 dH₂O and spun in a microfuge 5 minutes to pellet insoluble matter. 2µl of each cDNA-containing 15 supernatant were analyzed by electrophoresis on a 6% polyacrylamide gel. Autoradiography of the gel showed that the DNA in the pool of fractions 1-6 had an average size of 1100 base-pairs (bp) and included DNA in the 200 bp range and this pool was chosen for addition of "polyC 20 tails" to the 3' ends of the cDNA, using, generally, the homopolymeric tailing procedure described in McCandliss et al., page 601 et seq., supra. A 5000 molar excess of dCTP over 3' cDNA ends was found to give good results.

The reaction mixture was as follows:

20µl cDNA (about 43 ng)

- ³H dCTP (645 pmol, lyophilized)

2.4µl 10X TdT buffet*

1.6ul dH2 0

30 24.0µl

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*10X TdT buffer = 1.4M potassium cacodylate/0.3M Tris-HCl, pH 7.0/10mM CoCl₂/1mM DTT)

The reaction mixture was preincubated to 37°C for 2 minutes, 2µl were removed for use in calculations, then 2µl (6.66 units) P-L Biochemicals terminal deoxynucleo-

tidyl transferase were added and incubation at 37°C was continued for 5 minutes. Calculations based on incorporation of ³B dCTP indicated that the 3'ends of the cDNA now carried "polyC tails" an average of 14 nucleotides in length. 80µl T.E. buffer (10mM Tris-HCl, pB 7.6/1mM EDTA) were added to the DNA and the solution was extracted with an equal volume of (1:1) phenol:chloroform. The organic phase was then retracted with 100µl dB₂O and the two aqueous phases were combined.

The C-tailed double-stranded cDNA was then annealed to plasmid pBR322 DNA which had been linearized with the restriction endonuclease PstI, then "G-tailed" by the homopolymeric tailing method. The complementary single-stranded C and G "tails" will anneal, producing recombinant plasmids with cDNA inserts at the PstI site.

200µl cDNA, C-tailed (39.2 ng) 10.5µl pBR322-PstI, G-tailed (302 ng) 93µl 10X buffer*

20 <u>626.5</u> µ1 dH₂O 930µ1

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The reaction mix was placed in an insulated water bath at 70°C. The bath was then transferred to a 37°C room and allowed to cool slowly to 37°C overnight, then transferred to room temperature, where the bath cooled to 30°C over several hours. The reaction mixture was then stored at 4°C.

*(10X annealing buffer = 1.5M NaCl/100mM Tris-HCl, pH7.5/10mM EDTA)

E. coli HB101 cells were made competent for transformation by known calcium chloride treatment procedures. 200µl aliquots of competent HB101 cells were each combined with 40µl of the annealing reaction mixture and kept on ice 20 minutes, then heat-shocked at 42°C for 2 minutes. 2.8 mls Luria broth were added to

each tube and incubated at 37°C for 1 hour. The tubes' contents were aliquoted (1/2 ml aliquots) into tubes containing Luria broth plus 0.7% agar, and then were poured onto Luria broth-agar plates containing 25µg/ml tetracycline and incubated at 37°C until colonies appeared.

Only those cells transformed by pBR322 (with or without a cDNA insert) can grow on tetracycline plates. Approximately 2500 transformant colonies grew on the plates.

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EXAMPLE III Isolation of a Full-Length HSA cDNA

The transformants were initially screened with a rat serum albumin (RSA) cDNA fragment. The RSA cDNA fragment was obtained from a pBR322 plasmid containing a 15 2000 bp RSA cDNA insert. This recombinant plasmid is similar to, but contains a longer cDNA insert than, the plasmid prAlbI described in Proc. Nat'l. Acad. Sci. USA, 76, 4370 (1979). A 1480 bp rat serum albumin (RSA) fragment was isolated by digesting the plasmid carrying 20 the RSA cDNA with the restriction endonuclease BstBII (all restriction endonucleases used in these examples were used according to manufacturer's specifications.) The fragment was then radioactively labeled with a32P by the "nick translation" procedure (Maniatis et al. PNAS 25 USA, 72:3961 (1975)).

About 80 10-ml cultures of individual transformants were grown and plasmid DNA was isolated by known plasmid "mini-prep" procedures. The partially purified plasmid DNAs were subjected to electrophoresis on 0.8% agarose gels. The DNA was transferred from the gels to nitrocellulose filters using the "Southern blotting"

technique (Southern, E.M. <u>J. Molec. Biology</u> <u>98</u>, 503 (1975)).

The nitrocellulose filters were immersed for 2 hours at 42°C in prehybridization solution (50% formamide/5X SSC*/0.05M NaPO,, pH 6.5/5X Denhardt's*/100µg/ml salmon sperm DNA). The filters were then transferred into hybridization solution (50% formamide/10% dextran sulfate/5X SSC/20mM NaPO,, pH 6.5/1X Denhardt's/50µg/ml salmon sperm DNA.) The nick-translated 1480bp RSA fragment prepared above was heated at 100°C for 5 minutes, then quick cooled on ice, and this probe was added to the hybridization solution at 2 X 10⁵ cpm probe per ml of solution. The filters were incubated in the hybridization solution at 42°C for 18 hours, then washed twice in 2XSSC and once in 0.1X SSC at room temperature.

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Autoradiography of the filters revealed non-specific hybridization of the probe to all plasmid DNAs. Therefore, several Southern blot filters were washed in 2XSSC at various temperatures from 65°C to 80°C. DNA from one plasmid on a filter washed at 65°C hybridized strongly with the probe.

DNA sequencing revealed that the "positive" clone, called 6C3, was a partial-length human serum albumin clone. Plasmid DNA was isolated from a culture of 6C3 and digested with the restriction endonuclease PstI. One of the resulting HSA cDNA fragments, about 475bp in length, was isolated and "nick translated" for use as a

^{30 *50%} Denhardt's stock = 1% polyvinylpyrrolidone/1% ficoll/1% bovine serum albumin.

¹XSSC = 150mM NaCl/15mM sodium citrate, pH 6.8 with citric acid

probe. The entire bank of approximately 2500 clones was screened with this probe using a modification of the hybridization procedure of Grunstein et al., supra.

The transformant colonies were individually picked 5 from the plates into separate wells in 96-well microtiter plates containing Luria broth plus 0.2% glucose plus 25µg/ml tetracycline and incubated at 37°C ovenight. Using a transfer device with 48 metal prongs, samples of each culture were transferred to two Luria 10 broth/agar/tetracycline plates, one plate previously overlaid with a nitrocellulose filter, and incubated at 37°C 2 days. The filters were then placed successively on Whatman filter paper soaked in one of the following solutions: 0.5M NaOH; 1MTris, pH7.4; 1M Tris, pH7.4; 15 2XSSC; 90% ethanol, and 90% ethanol (in that order, 7 minutes per solution.) The nitrocellulose filters were then baked in vacuo at 80°C for 2 hours.

Prehybridization and hybridization procedures were as described above, except that the three washes were at room temperature. 90 positive hybridization signals were detected by autoradiography. Some of the "positive clones" were further analyzed by restriction analysis (e.g. PstI digestion) and hybridization of "Southern blots" as above.

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A clone bearing a full length HSA cDNA was identified and confirmed by DNA sequencing. The recombinant plasmid containing this HSA cDNA insert was termed pGX401 and is shown in figure 4. A partial restriction map of the HSA cDNA is shown in Figure 1, while Figure 2 shows the DNA sequence (5'+3' strand) of the cloned gene and the amino acid sequence it specifies.

A sample of \underline{E} . \underline{coli} HB101 transformed with pGX401 has been deposited at the U.S. Dept. of Agriculture

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Northern Regional Research Center in Peoria, Illinois. under accession number NRRL B-15784.

EXAMPLE IV

DNA Sequence Analysis of HSA cDNA Prepared from Buman Liver Samples Taken from Different Individuals

In comparing the DNA sequence of the HSA cDNA insert in pGX401 (Example III) with the cDNA sequences published by Lawn et al., supra, and Dugaiczyk et al., supra, two codon differences were found that predict amino acid differences. The pGX401 sequence and the sequence reported by Lawn et al. indicated that codon 97 of the mature protein was GAG (glutamic acid), while Dugaiczyk et al. reported it to be GGG (glycine). In the pGX401 sequence and the sequence reported by Dugaiczyk codon 396 also was reported to be GAG (glutamic acid), and Lawn et al. reported that codon to be AAG (lysine).

To gain some insight into whether these differences represented true protein polymorphisms or merely experimental artifacts, the DNA sequence in the regions of codons 97 and 396 was determined for several new independent HSA genes.

Messenger RNA (mRNA) was isolated from normal human liver tissue taken from four different individuals. The procedures of Example I were followed except that sucrose gradient fractionation of oligo (dT)-cellulose-purified mRNA was omitted. Double stranded cDNA was synthesized from this mRNA template by the procedures described in Example II and poly(dC) "tails" were added according to Deng and Wu, NAR 9:4123, 1981.

The vector into which the dC-tailed cDNA was inserted was plasmid pGX1066. This plasmid comprises the phage λtR_1 transcription terminator upstream of a bank of ten closely-spaced unique restriction sites, which in turn is upstream of the $\lambda 4S$ transcription terminator.

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E. coli strain GX1170 [F' leu hsdR thi sup8 gal-1,2 lac xyl ara trpC9830 lacIq] transformed with pGX1066 has been deposited with the American Type Culture Collection, Rockville, Maryland, as ATCC No. 39955.

Plasmid pGX1066 was linearized with PstI and poly(dG) tails were added using the homopolymeric tailing method described by Deng and Wu (Nucleic Acids Res., 9: 4173 (1981)). The vector DNA and cDNA were then annealed as described in Example II. E. coli strain DH1 cells

(F-, endA1, hsdR17 (R_k-, M_k-), supE44, thi1, λ-, recA1, gyrA96, relA1] were made competent and transformed with the annealing reaction mix. Both E. coli strain DH1 and the transformation procedure used are described by D. Hanahan (J. Molec. Biol., 166: 557 (1983)). Transfor-

mants were plated on LM plates (1% (w/v) Bacto tryptone, 0.5% (w/v) yeast extract, 10mM NaCl, 10mM MgSO4.7H2O, 1.5% (w/v) Bacto agar) with 35µg/ml ampicillin added.

Transformed E. coli colonies were screened for the presence of HSA sequences by Grunstein-Hogness filter hybridization (Gergen et al., 1979, Nuc. Acids. Res. 7:2115; Wallace et al., 1981, Nuc. Acids Res. 9:879) using kinased oligomers or nick-translated HSA cDNA fragments as probes. For identification of clones carrying HSA cDNA containing codon 396, a synthetic oligonucleotide, 5' TTGTACTCTCCAAGCTGC 3', corresponding to codons 397-402 (and the last nucleotide of codon 396) was used. For detection of clones carrying HSA cDNA

oligonucleotides, 5' TCTCTTCATTGTCATGAAAAGC 3',

corresponding to codons 126-132 (and one nucleotide of
codon 133), or 5' TTCTTGTTTTGCACAGC 3', corresponding to
codons 90 (last 2 nucleotides) - 95, or a nick-translated
HSA fragment (derived from pGX401), corresponding to
codons -1 to 364 was used. Upon identification of clones
containing the HSA sequence of interest, restriction

containing codon 97, either of two synthetic

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fragments were subcloned into an M13 phage. BSA cDNA-carrying phage were identified by screening plaques according to the procedure of Benton and Davis (Science, 196:180 (1977)). The DNA sequence was determined with these M13 clones by the dideoxy method (Biggin et al., Proc. Nat. Acad. Sci., U.S.A. 80:3963 (1983)).

By the procedures described above, transformants containing HSA cDNA that included codon 396 were derived from all four human livers. Transformants containing HSA cDNA that included codon 97 were derived from only two of the four livers. The DNA sequence in all cases (including 60 to 100 base pairs on each side of the codon in question) matched the sequence determined for pGX401.

Messenger RNA then was isolated from normal human liver samples taken from two more individuals, and the sequence at codon 97 was determined using a modification of the Sanger sequencing procedure in which reverse transcriptase was used to copy the single-stranded RNA template. A synthetic oligonucleotide,

template. A synthetic oligonucleotide,
5' TGTCTCTTCATTGTCATGAAAAGC 3', corresponding to codons
126-133, was used as a primer. The mRNA, purified by
oligo (dT)-cellulose chromatography as previously
described, was incubated in a reaction volume of 2μl
containing 10 mM Tris · HCl (pH 8.3), 140 mM KCl, 10 mM
MgCl₂, 20 mM β-mercaptoethanol, 1.6 mM dNTP, 0.2 mM
ddNTP, 250 ng RNA, 5 ng kinased primer and 1.88 units
reverse transcriptase (Life Sciences, Inc.). After
overlaying the solution with 4 μl of mineral oil the
reaction was incubated at 42°C for fifteen minutes and
was terminated by the addition of 7 μl of 250 mM Na₂
EDTA. The mineral oil was extracted with ether and
removed with a drawn-out pasteur pipette. Formamide
loading buffer was added to the samples prior to electrophoresis on a urea sequencing gel. The gels were run

until the bromphenol blue tracking dye had migrated to

the bottom. They then were dried under vacuum and exposed to X-ray film with two intensifying screens for periods between twelve hours and several days.

The HSA sequence at codon 97 for both liver samples was identical to the sequence at codon 97 in pGX401. (See Figure 5.) The reliability of the technique to determine nucleotide sequence from mRNA was evaluated using polyA+ RNA prepared from the liver that was the source of the cDNA originally cloned in pGX401. The results (Figure 5) showed that the sequence determined in 10 this manner was identical to the sequence originally determined in pGX401.

CLAIMS FOR THE DESIGNATED STATES: BE, DE, FR, IT, LU, NL, SE, CH and UK

- 1. A synthetic gene coding for human serum albumin.
- 2. An isolated human serum albumin gene.
- 3. An isolated prepro-human serum albumin gene.
- 4. A human serum albumin gene as claimed in claim 1, comprising the following deoxyribonucleotide sequence which corresponds to the indicated amino acid sequence:

Bis Ser Glu Val Ala Lys GAY GCX CAY AAM QRS GAM GTX GCX Gly Arg Phe Leu Lys Asp CAY LGN TTY AAM GAY YTZ GGX GAM Va1 Leu Ala Leu Glu Asn Phe Lys GAM AAY TTY AAM GCX YTZ GTX YTZ Phe Ala GlnTyr Leu ATH GCX TTY GCX CAM TAY YTZ Phe Glu Asp His Val Cys Pro CAM TGY CCX TTY GAM GAY CAY GTX Val Asn Glu Val -Thr Lys Leu AAM YTZ GTX AAY GAM GTX ACX GAM Ala Lys Thr Cys Val Ala Asp TTY GCX AAM ACX TGY GTX GCX GAY Cys Asp Glu Asn Ser Ala GAM QRS GCX GAM AAY TGY GAY AAM Thr Leu Phe Gly Asp Leu His Ser QRS YTZ CAY ACX YTZ TTY GGX GAY Thr Leu Cys Thr Val Ala Leu AAM Y.T Z TGY ACX GTX GCX ACX YTZ Met Gl y Gl u Arg Glu Thr Tyr LGN GAM ACX TAY GGX GAM ATG GCX Glu GlnCys Ala Lys Cys GAY TGY TGY GCX AAM CAM GAM CCX

Glu Arg Asn Glu Cys Phe Leu GAM LGN AAY GAM TGY TTY YTZ CAM Lys Asp Asp Asn Pro Asn Leu CAY AAM GAY GAY AAY CCX AAY YTZ Arg Leu Val Arg Pro Glu Val CCX LGN YTZ GTX LGN CCX GAM GTX Val Met Cys Thr Phe Ala Ris GAY GTX ATG TGY ACX GCX TTY CAY Asp Asn Glu Glu Thr Phe Lvs GAY AAY GAM GAM ACX TTY YTZ AAM Lys Tyr Leu Ile Tyr Glu AAM TAY YTZ TAY GAM ATH GCX LGN His Pro Tyr Phe Thr Ala LGN CAY CCX TAY TTY ACX GCX CCX Leu Leu Phe Ala Phe Lys GAM YTZ YTZ TTY TTY GCX AAM LGN Lys Ala Ala Phe Thr TAY AAM GCX GCX TTY ACX GAM TGY Gln Cys Ala Ala Asp Lys Ala TGY GCX CAM GCX GAY AAM GCX GCX Cys Leu Phe Pro Lys Leu Asp TGY YTZ TTY CCX AAM YTZ GAY GAM Arg Gl y Glu Asp Lys Ala YTZ LGN GAY GAM GGX AAM GCX QRS Ser Ala Lys Gln Arg Leu Lys Cys QRS GCX AAM CAM LGN YTZ AAM TGY Ala Ser Leu Gln Phe Gl y Lys Glu GCX QRS YTZ CAM AAM TTY GGX GAM Ala Arq Phe Lys Ala Trp Ala Val L G N G C X T T Y A A M G C X T G G G C X G T X Ala Arg Leu Ser Gln Arg Phe GCX LGN YTZ QRS CAM LGN TTY CCX Lys Ala Glu Phe Ala Gl u Val AAM GCX GAM TTY GCX GAM GTX QRS Lys Phe Asp Val Thr Leu Thr Lys AAM TTY GTX ACX GAY YTZ ACX AAM Val His Glu Thr Cys Cys GTX CAY ACX GAM TGY TGY CAY GGX

30 Asp Leu Leu Glu Cys Ala Asp GAY YTZ YTZ GAM TGY GCX GAY GAY Ala Asp Leu Ala Lys Tyr L GN GCX GAY YTZ GCX AAM TAY ATH Glu Asn Cys Gln Asp Ser Ile Ser TGY GAM AAY CAM GAY QRS ATH QRS Ser Lys Leu Lys Glu Cys Cys Glu QRS AAM YTZ AAM GAM TGY TGY GAM Leu Phe Lys Pro Glu Lys Ser Bis AAM CCX YTZ TTY GAM AAM QRS CAY Ile Ala Glu Val Glu neA TGY ATH GCX GAM GTX GAM AAY GAY Met Pro Ala Asp Phe Pro Ser GAM ATG CCX GCX GAY TTY CCX QRS Phe Ala Val Asp Phe Val Glu Ser T T Y G C X G T X G A Y T T Y G T X G A M Q R S Cys Lys Asp Val Tyr Asn AAM GAY GTX TGY AAM AAY TAY GCX Ala Lys Val Phe Leu Asp GAM GCX AAM GAY GTX TTY YTZ GGX Met Phe Phe Tyr Glu Tyr Ala Arg A T G T T Y T T Y T A Y G A M T A Y G C X L G N Met His Pro Val Asp Tyr Ser LGN CAY CCX GAY TAY QRS: GTX GTX Leu Leu Arg Leu Ala Lys Thr Y T Z Y T Z Y T Z L G N Y T Z G C X A A M A C X Lys Glu Thr Thr Leu Glu Lys Tvr TAY GAM ACX ACX YTZ GAM AAM TGY Ala Ala Pro His Ala Asp TGY GCX GCX GCX GAY CCX CAY GAM Phe Ťyr Ala Lys Val Asp TGY TAY GCX AAM GTX TTY GAY GAM Val Glu Glu Pro Lys Pro Pro TTY AAM CCX CCX GTX GAM GAM CCX

Ile Gln Phe Lys Gln Asn Asn CAM AAY TTY ATH AAM CAM AAY TGY Glu Gln Gly Leu Phe Leu GAM YTT TTY GAM CAM YTT GGX GAM Lys Phe Gln Asn Ala Leu TAY AAM TTY CAM AAY GCX YTZ TTY Tyr Thr Lys Lys Val Arg GTX LGN TAY ACX AAM AAM GTX CCX Thr Leu Ser Thr Pro Leu Val CAM YTZ QRS ACX CCX ACX YTZ GTX Asn Gly Glu Val Ser Arg Leu GAM GTX ORS LGN AAY YTZ GGX AAM Gly Val Ser Lys Cys Cys Lys GTX GGX Q 9 S AAM TGY TGY AAM CAY Glu Ala Lys Arg Met Pro CCX GAM GCX AAM LGN ATG CCX TGY Ala Glu Asp Tyr Leu Ser Val Val G C X G A M G A Y T A Y Y T Z Q R S G T X G T X Gln Val Leu Leu Cys Leu Asn YTZ AAY CAM YTZ TGY GTX YTZ CAY Lys Thr Pro Val Ser Asp GAM AAM ACX CCX GTX QRS GAY LGN Thr Cys Cys Glu Ser Thr Lys GTX ACX AAM TGY TGY ACX GAM QRS Gly Val Asn Arg Arg Pro YTZ GTX AAY LGN LGN CCX GGX TTY Glu Val Glu Ala Leu Asp Thr QRS GCX YTZ GAM GTX GAY GAM ACX Tyr Val Pro Lys Glu Phe Asn Ala TAYGTX CCX AAM GAM TTY AAYGCX Thr Phe His Ala Phe Thr GAM ACX TTY ACX TTY CAY GCX GAY Ser Cys Thr Leu Glu Lys ATH TGY ACX YTZ QRS GAM AAM GAM

32 Gln Arg Ile Lys Glu Lys Thr LGN CAM ATH AAM AAM GAM ACX GCX Leu Val Glu Leu Val Lys His Lys YTZ GTX GAM YTZ GTX AAM CAY AAM Lys Ala Thr Lys Glu Glu Leu CCX AAM GCX ACX AAM GAM GAM YTZ Lys Ala Val Met Asp Asp AAM GCX GTX ATG GAY GAY TTY GCX Val Phe Ala Glu Lys Cys Cys Lys GCX TTY GTX GAM AAM TGY TGY AAM Asp Glu Asp Lys Thr Cys Phe GCX GAY GAY AAM GAM ACX TGY TTY Glu Ala Glu Gly Val Lys Lys Leu GCX GAM GAM GGX AAM AAM YTZ GTX Glu Ala Ser Ala Val Leu Gly GCX GCX QRS GAM GCX GTX YTZ GGX Leu Y T Z T A A

wherein, the 5' to 3' strand, beginning with the amino terminus and the amino acids for which each triplet codes are shown, and wherein the abbreviations have the following standard meanings:

A is deoxyadenyl

T is thymidyl

G is deoxyguanyl

C is deoxycytosyl

X is A, T, C or G

Y is T or C

When Y is C, Z is A, T, C or G

When Y is T, Z is A or G

H is A, T or C

Q is T or A

When Q is T, R is C and S is A, T, C or G

When Q is A, R is G and S is T or C

M is A or G L is A or C When L is A, N is A or G When L is C, N is A, T, C or G GLY is glycine ALA is alanine VAL is valine LEU is leucine ILE is isoleucine SER is serine THR is threonine PHE is phenylalanine TYR is tyrosine TRP is tyryptophan CYS is cysteine MET is methionine ASP is aspartic acid GLU is glutamic acid LYS is lysine ARG is arginine HIS is histidine PRO is proline

GLN is glutamine ASN is asparagine

5. A prepro-serum albumin gene as claimed in claim 1 comprising the following deoxyribonucleotide sequence:

Trp Val Thr Met Lys ATG AAM TGG GTX ACX T T Y Phe Ser Leu Leu Leu ATH QRS YTZ YTZ TTYYZ Arg Ser Ala Tyr Ser QRS Q'RS GCX TAY QRS LGN Phe Arg Asp Ala His Ly s Val Arg GTX TTY LGN LGN GAY GCX CAY AAM

34 Arg Phe His Lys Glu Val Ala QRS GAM GTX GCX CAY LGN TTY AAM Asp Leu Gly Glu Glu Asn Phe Lys G A Y Y T Z G G X G A M G A M A A Y T T Y A A M Leu Ile Phe Val Ala Leu GCX YTZ GTX YTZ ATH GCX TTY GCX Gln Gln Cys Pro Tyr Leu CAM TAY YTZ CAM CAM TGY CCX TTY Leu Val Asn Asp His Val Lys GAM GAY CAY GTX AAM YTZ GTX AAY Glu Lys Phe Ala Val Thr GAM GTX ACX GAM TTY GCX AAM ACX Ala Ser Val Ala Asp Glu TGY GTX GCX GAY GAM QRS GCX GAM Asn Cys. Asp Lys Ser Leu His Thr AAYTGYGAYAAMQRSYTZCAYACX Leu Phe Gly Asp Lys Leu Cys Thr Y T Z T T Y G G X G A Y A A M Y T Z T G Y A C X Val Ala Thr Leu Arg Glu Thr Tyr G T X G C X A C X Y T Z L G N G A M A C X T A Y Gly Glu Met Ala Asp Cys Cys Ala G G X G A M A T G G C X G A Y T G Y T G Y G C X Asn Pro Glu Arg Gln Glu Lvs AAM CAM GAM CCX GAM LGN AAY GAM Lys Asp His Gln Phe Leu TGY TTY YTZ CAM CAY AAM GAY GAY Asn Pro Asn Leu Pro Arg Leu Val A A Y C C X A A Y Y T Z C C X L G N Y T Z G T X Arg Pro Glu Val Asp Val Met Cys LGN CCX GAM GTX GAY GTX ATG TGY Glu His Asn Phe qaA Ala ACX GCX TTY CAY GÂY AAY GAM GAM Leu Lys Tyr Lys Phe Leu ACX TTY YTZ AAM AAM TAY YTZ TAY

35 Ile His Ala Arg Arg Pro GAM ATH GCX LGN LGN CAY CCX TÂY Thr Ala Pro Glu Leu Leu TTY ACX GCX CCX GAM YTZ YTZ TTY Ala Lys Arg Tyr Lys Ala TTY GCX AAM LGN TAY AAN GCX GCX Cys Cys Ala Gln Thr Glu TTY ACX GAM TGY TGY GCX CAM GCX Lys Ala Ala Cys Leu Phe GAY AAM GCX GCX TGY YTZ TTY CCX Lys Leu Asp Glu Leu Arg Asp Glu A A M Y T Z G A Y G A M Y T Z L G N G A Y G A M Arg Lys Ala Ser Ser Ala Lys G G X A A M G C X Q R S Q R S G C X A A M C A M Cys Ala Ser Leu Leu Lys LGN YTZ AAM TGY GCX QRS YTZ CAM Gly Phe Ala Phe Glu Arg Lvs AAM TTY GGX GAM LGN GCX TTY AAM Ala Leu Ser Ala Val Arg Trp GCX TGG GCX GTX GCX LGN YTZ QRS Pro Lys Ala Glu Phe Arg Phe CAM LGN TTY CCX AAM GCX GAM TTY Val Thr Phe Glu Lys Val Ser GCX GAM GTX QRS AAM TTY GTX ACX Th r Thr Lys Val His Leu GAY YTZ ACX AAM GTX CAY ACX GAM Gly Asp Leu Leu Cys His TGY TGY CAY GGX GAY YTZ YTZ GAM Ala Asp Asp Leu Asp Arg TGY GCX GAY GAY LGN GCX GAY YTZ Ile Cys Glu Lys Tyr GCX AAM TAY ATH TGY GAM AAY CAM Ile Ser Lys Leu Ser Ser GAY QRS ATH QRS QRS AAM YTZ AAM

36 Glu Cys Cys Glu Lys Pro Leu Phe G A M T G Y T G Y G A M A A M C C X Y T Z T T Y Ser His Cys Ile Ala Lys GAM AAM QRS CAY TGY ATH GCX GAM Val Glu Asn Asp Glu Met Pro Ala G T X G A M A A Y G A Y G A M A T G C C X G C X Pro Val Phe Ala Phe Ser GAY TTY CCX QRS TTY GCX GTX GAY дaя Glu Val Val Ser Lys Cys TTY GTX GAM QRS AAM GAY GTX TGY Tyr Glu Lys Ala Ala Asp Lys Asn AAM AAY TAY GCX GAM GCX AAM GAY Gly Met Phe Phe Tyr Leu G T X T T Y Y T Z G G X A T G T T Y T T Y T A Y Glu Tyr Ala Arg Arg His Pro Asp G A M T A Y G C X L G N L G N C A Y C C X G A Y Val Leu Leu Leu Tyr Ser Val TAY QRS GTX GTX YTZ YTZ LGN Gl u Lys Tyr Thr Thr Ala YTZ GCX AAM ACX TAY GAM ACX ACX Ala Cys Ala Leu Glu Lvs Cys YTZ GAM AAM TGY TGY GCX GCX Ris Gl u Cys Tyr Ala Asp Pro GAY CCX CAY GAM TGY TAY GCX AAM Asp Glu Phe Lys Pro Phe G T X T T Y G A Y G A M T T Y A A M C C X C C X Pro Gln Asn Phe Glu Glu GTX GAM GAM CCX CAM AAY TTY ATH Glu Phe asa Cys Leu Gln AAM CAM AAY TGY GAM YTZ TTY GAM Gln Leu Gly Glu Tyr Lys Phe Gln CAMYTZGGXGAMTAYAAMTTYCAM Tyr Leu Phe **Val** Arg Ala AAY GCX YTZ TTY GTX LGN TAY ACX

Gln Leu Ser Thr Pro Val Lys AAM AAM GTX CCX CAM YTZ QRS ACX Lys Ser Arg Glu Val Leu Va1 Thr CCX ACX YTZ GTX GAM GTX QRS LGN Asn Leu Gly Lys Val Gly Ser Lys AAYYTZGGXAAMGTXGGXQRSAAM Ala Glu His Pro Lys Cys Cvs TGY TGY AAM CAY CCX GAM GCX AAM Arg Met Pro Cys Ala Glu Asp Tyr L G N A T G C C X T G Y G C X G A M G A Y T A Y Leu Leu Gln Val Asn Val YTZ QRS GTX GTX YTZ AAY CAM YTZ Lys Thr Leu His Glu Val Суз TGY GTX YTZ CAY GAM AAM ACX CCX Arg Val Th r Lys Ser Asp Val GTX QRS GAY LGN GTX ACX AAM TGY Ser Leu Val Asn Arq Thr Glu TGY ACX GAM QRS YTZ GTX AAY LGN Arg Pro Gly Phe Ser Ala Leu Glu L G N C C X G G X T T Y Q R S G C X Y T Z G A M Lys Val Asp Glu Thr Tyr Val Pro Lys G T X G A Y G A M A C X T A Y G T X C C X A A M Tyr Val Thr Phe Glu Ala Asn Phe GAM TTY AAY GCX GAM ACX TTY ACX Glu Thr Leu Cys Ile Ala Asp His TTY CAY GCX GAY ATH TGY ACX YTZ Ile Lys Arg Gl n Glu Lys Glu Ser QRS GAM AAM GAM LGN CAM ATH AAM Gl u Val Leu Glu Thr Ala AAM GAM ACX GCX YTZ GTX GAM YTZ Pro Lys Ala Lys Lys His GTX AAM CAY AAM CCX AAM GCX ACX Ala Va l Leu Lys Glu Glu Lvs AAM GAM GAM YTZ AAM GCX GTX ATG Phe Va l Ala Ala Phe Asp GAY GAY TTY GCX GCX TTY GTX GAM

Cys Cys Lys Ala Asp Asp AĀM TĒY TĒY AĀM GCX GAY GAY AĀ M Ala Glu Glu Cys Phe Gl v Thr GAM ACX TGY TTY GCX GAM GAM GGX Ser Ala Ala Lys Leu Val AAM AAM YTZ GTX GCX GCX QRS GAM Gly Ala Val Leu Leu GCX GTX YTZ GGX YTZ TAA

wherein the 5' and 3' strand, beginning with the amino terminus, and the amino acids for which each triplet codes are shown, and wherein the abbreviations are defined as in claim 4.

A human serum albumin gene as claimed in claim 4 comprising the following deoxyribonucleotide sequence: GAT GCA CAC AAG AGT GAG GTT GCT CAT CGG TTT AAA GAT TTG GGA GAA GAA AAT TTC AAA GCC TTG GTG TTG ATT GCC TTT GCT CAG TAT CTT CAG CAG TGT CCA TTT GAA GAT CAT GTA AAA TTA GTG AAT GAA GTA ACT GAA TTT GCA AAA ACA TGT GTT GCT GAT GAG TCA GCT GAA AAT TGT GAC AAA TCA CTT CAT ACC CTT TTT GGA GAC AAA TTA TGC ACA GTT GCA ACT CTT CGT GAA ACC TAT GGT GAA ATG GCT GAC TGC TGT GCA AAA CAA GAA CCT GAG AGA AAT GAA TGC TTC TTG CAA CAC AAA GAT GAC AAC CCA AAC CTC CCC CGA TTG GTG AGA CCA GAG GTT GAT GTG ATG TGC ACT GCT TIT CAT GAC AAT GAA GAG ACA TIT TIG AAA AAA TAC TIA TAT GAA ATT GCC AGA AGA CAT CCT TAC TTT TAT GCC CCG GAA CTC CTT TTC TTT GCT AAA AGG TAT AAA GCT GCT TTT ACA GAA TGT TGC CAA GCT GCT GAT AAA GCT GCC TGC CTG TTG CCA AAG CTC GAT GAA CTT CGG GAT GAA GGG AAG GCT TCG TCT GCC AAA CAG AGA CTC AAG TGT GCC AGT CTC CAA AAA TTT GGA GAA AGA GCT TTC AAA GCA TGG GCG GTG GCT CGC CTG AGC CAG AGA TTT CCC AAA GCT GAG TTT GCA GAA GTT TCC AAG TTA GTG ACA GAT CTT ACC AAA GTC CAC ACG GAA TGC TGC CAT GGA GAT CTG CTT GAA TGT GCT GAT GAC AGG GCG GAC CTT GCC AAG TAT ATC TGT GAA AAT CAA GAT TCG ATC TCC AGT AAA CTG AAG GAA TGC TGT GAA AAA CCT CTG TTG GAA AAA TCC CAC TGC ATT GCC GAA GTG GAA AAT GAT GAG ATG CCT GCT GAC TTG CCT TCA TTA GCT GCT GAT TTT GTT GAA AGT AAG GAT GTT TGC AAA AAC TAT GCT GAG GCA AAG GAT GTC TTC CTG GGC ATG TTT TTG TAT GAA TAT GCA AGA AGG CAT CCT GAT TAC TCT GTC GTG CTG CTG CTG AGA CTT GCC AAG ACA TAT GAA ACC ACT CTA GAG AAG TGC TGT GCC GCT GCA GAT CCT CAT GAA TGC TAT GCC AAA GTG TTC GAT GAA TTT AAA CCT CCT GTG GAA GAG CCT CAG AAT TTA ATC AAA CAA AAT TGT GAG CTT TTT GAG CAG CTT GGA GAG TAC AAA TTC CAG AAT GCG CTA TTA GTT CGT TAC ACC AAG AAA GTA CCC CAA GTG TCA ACT CCA ACT CTT GTA GAG GTC TCA AGA AAC CTA GGA AAA GTG GGC AGC AAA TGT TGT AAA CAT CCT GAA GCA AAA AGA ATG CCC TGT GCA GAA GAC TAT CTA TCC GTG GTC CTG AAG CAG TTA TGT GTG TTG CAT GAG AAA ACG CCA GTA AGT GAC AGA GTC ACC AAA TGC TGC ACA GAA TCC TTG GTG AAC AGG CGA CCA 'IGC TTT TCA GCT CTG GAA GTC GAT GAA ACA TAC GTT CCC AAA GAG TTT AAT GCT GAA ACA TTC ACC TTC CAT GCA GAT ATA TGC ACA CTT TCT GAG AAG GAG AGA CAA ATC AAG AAA CAA ACT GCA CTT GTT GAG CTC GTG AAA CAC AAG CCC AAG GCA ACA AAA GAG CAA CTG AAA GCT GTT ATG GAT GAT TTC GCA GCT TTT GTA GAG AAG TGC TGC AAG GCT GAC GAT AAG GAG ACC TGC TTT GCC GAG GAG GGT AAA AAA CTT GTT GCT GCA AGT CAA GCT GCC TTA GGC TTA TAA wherein the 5' to 3' strand, beginning with the amino terminus is shown, and wherein the abbreviations are defined as in claim 4.

7. A human prepro-serum albumin gene as claimed in claim 5 comprising the following deoxyribonucleotide sequence: ATG AAG TGG GTA ACC TTT ATT TCC CTT CTT TTT CTC TTT AGC TCG GCT TAT TCC AGG GGT GTG TTT CGT CGA GAT GCA CAC AAG AGT GAG GTT GCT CAT CGG TTT AAA GAT TTG GGA GAA GAA AAT TTC AAA GCC TTG GTG TTG ATT GCC TTT GCT CAG TAT CTT CAG CAG TGT CCA TTT GAA GAT CAT GTA AAA TTA GTG AAT GAA GTA ACT GAA TTT GCA AAA ACA TGT GTT GCT GAT GAG TCA GCT GAA

AAT TGT GAC AAA TCA CTT CAT ACC CTT TTT GGA GAC AAA TTA TGC ACA GTT GCA ACT CTT CGT GAA ACC TAT GGT GAA ATG GCT GAC TGC TGT GCA AAA CAA GAA CCT GAG AGA AAT GAA TGC TTC TTG CAA CAC AAA GAT GAC AAC CCA AAC CTC CCC CGA TTG GTG AGA CCA GAG GTT GAT GTG ATG TGC ACT GCT TTT CAT GAC AAT GAA GAG ACA TIT ITG AAA AAA TAC ITA TAI GAA AII GCC AGA AGA CAT CCT TAC TTT TAT GCC CCG GAA CTC CTT TTC TTT GCT AAA AGG TAT AAA GCT GCT TTT ACA GAA TGT TGC CAA GCT GCT GAT AAA GCT GCC TGC CTG TTG CCA AAG CTC GAT GAA CTT CGG GAT GAA GGG AAG GCT TCG TCT GCC AAA CAG AGA CTC AAG TGT GCC AGT CTC CAA AAA TTT GGA GAA AGA GCT TTC AAA GCA TGG GCG GTG GCT CGC CTG AGC CAG AGA TTT CCC AAA GCT GAG TTT GCA GAA GTT TCC AAG TTA GTG ACA GAT CTT ACC AAA GTC CAC ACG GAA TGC TGC CAT GGA GAT CTG CTT GAA TGT GCT GAT GAC AGG GCG GAC CTT GCC AAG TAT ATC TGT GAA AAT CAA GAT TCG ATC TCC AGT AAA CTG AAG GAA TGC TGT GAA AAA CCT CTG TTG GAA AAA TCC CAC TGC ATT GCC GAA GTG GAA AAT GAT GAG ATG CCT GCT GAC TTG CCT TCA TTA GCT GCT GAT TTT GTT GAA AGT AAG GAT GTT TGC AAA AAC TAT GCT GAG GCA AAG GAT GTC TTC CTG GGC ATG TTT TTG TAT GAA TAT GCA AGA AGG CAT CCT GAT TAC TCT GTC GTG CTG CTG AGA CTT GCC AAG ACA TAT GAA ACC ACT CTA GAG AAG TGC TGT GCC GCT GCA GAT CCT CAT GAA TGC TAT GCC AAA GTG TTC GAT GAA TTT AAA CCT CCT GTG GAA GAG CCT CAG AAT TTA ATC AAA CAA AAT TGT GAG CTT TTT GAG CAG CTT GGA GAG TAC AAA TTC CAG AAT GCG CTA TTA GTT CGT TAC ACC AAG AAA GTA CCC CAA GTG TCA ACT CCA ACT CTT GTA GAG GTC TCA AGA AAC CTA GGA AAA GTG GGC AGC AAA TGT TGT AAA CAT CCT GAA GCA AAA AGA ATG CCC TGT GCA GAA GAC TAT CTA TCC GTG GTC CTG AAC CAG TTA TGT GTG TTG CAT GAG AAA ACG CCA GTA AGT GAC AGA GTC ACC AAA TGC TGC ACA GAA TCC TTG GTG AAC AGG CGA CCA TGC TTT TCA GCT CTG GAA GTC GAT GAA ACA TAC GTT CCC AAA GAG TTT AAT GCT GAA ACA TTC ACC TTC CAT GCA GAT ATA TGC ACA CTT TCT GAG AAG GAG AGA CAA ATC AAG AAA CAA ACT GCA CTT GTT GAG CTC GTG AAA CAC AAG CCC AAG GCA ACA AAA GAG CAA CTG AAA GCT GTT ATG GAT GAT TTC GCA GCT TTT GTA GAG AAG TGC TGC AAG GCT GAC GAT AAG GAG ACC TGC TTT GCC GAG GAG GGT AAA AAA CTT GTT GCT GCA AGT CAA GCT GCC TTA GGC TTA TAA wherein the 5' to 3' strand, beginning with the amino terminus is shown, and wherein the abbreviations are defined as in claim 4.

8. A human prepro-serum albumin gene as claimed in claim 7 comprised in the following deoxyribonucleotide sequence: 5'

TCTCTTCTGTCAACCCCACGCCTTTGGCACA ATG AAG TGG GTA ACC TTT ATT TCC CTT CTT TTT CTC TTT AGC TCG GCT TAT TCC AGG GGT GTG TTT CGT CGA GAT GCA CAC AAG AGT GAG GTT GCT CAT CGG TTT AAA GAT TTG GGA GAA GAA AAT TTC AAA GCC TTG GTG TTG ATT GCC TTT GCT CAG TAT CTT CAG CAG TGT CCA TTT GAA GAT CAT GTA AAA TTA GTC AAT GAA GTA ACT GAA TTT GCA AAA ACA TGT GTT GCT GAT GAG TCA GCT GAA AAT TGT GAC AAA TCA CTT CAT ACC CTT TTT GGA GAC AAA TTA TGC ACA GTT GCA ACT CTT CGT GAA ACC TAT GGT GAA ATG GCT GAC TGC TGT GCA AAA CAA GAA CCT GAG AGA AAT GAA TGC TTC TTG CAA CAC AAA GAT GAC AAC CCA AAC CTC CCC CGA TTG GTG AGA CCA GAG GTT GAT GTG ATG TGC ACT GCT TTT CAT GAC AAT GAA GAG ACA TTT TTG AAA AAA TAC TTA TAT GAA ATT GCC AGA AGA CAT CCT TAC TTT TAT GCC CCG GAA CTC CTT TTC TTT GCT AAA AGG TAT AAA GCT GCT TT ACA GAA TGT TGC CAA GCT GCT GAT AAA GCT GCC TGC CTG TTG CCA AAG CTC GAT GAA CTT CGG GAT GAA GGG AAG GCT TCG TCT GCC AAA CAG AGA CTC AAG TGT GCC AGT CTC CAA AAA TTT GGA GAA AGA GCT TTC AAA GCA TGG GCG GTG GCT CGC CTG AGC CAG AGA TTT CCC AAA GCT GAG TTT GCA GAA GTT TCC AAG TTA GTG ACA GAT CTT ACC AAA GTC CAC ACG GAA TGC TGC CAT GGA GAT CTG CTT GAA TGT GCT GAT GAC AGG GCG GAC CTT GCC AAG TAT ATC TGT GAA AAT CAA GAT TCG ATC TCC AGT AAA CTG AAG GAA TGC TGT GAA AAA CCT CTG TTG GAA AAA TCC CAC TGC ATT GCC GAA GTG GAA AAT GAT GAG ATG CCT GCT GAC TTG CCT TCA TTA GCT GCT GAT TTT GTT GAA AGT AAG GAT GTT TGC AAA AAC TAT GCT GAG GCA AAG GAT GTC TTC CTG GGC ATG TTT

TTG TAT GAA TAT GCA AGA AGG CAT CCT GAT TAC TCT GTC GTG CTG CTG CTG AGA CTT GCC AAG ACA TAT GAA ACC ACT CTA GAG AAG TGC TGT GCC GCT GCA GAT CCT CAT GAA TGC TAT GCC AAA GTG TTC GAT GAA TTT AAA CCT CCT GTG GAA GAG CCT CAG AAT TTA ATC AAA CAA AAT TGT GAG CTT TTT GAG CAG CTT GGA GAG TAC AAA TTC CAG AAT GCG CTA TTA GTT CGT TAC ACC AAG AAA GTA CCC CAA GTG TCA ACT CCA ACT CTT GTA GAG GTC TCA AGA AAC CTA GGA AAA GTG GGC AGC AAA TGT TGT AAA CAT CCT GAA GCA AAA AGA ATG CCC TGT GCA GAA GAC TAT CTA TCC GTG GTC CTG AAC CAG TTA TGT GTG TTG CAT GAG AAA ACG CCA GTA AGT GAC AGA GTC ACC AAA TGC TGC ACA GAA TCC TTG GTG AAC AGG CGA CCA TGC TTT TCA GCT CTG GAA GTC GAT GAA ACA TAC GTT CCC AAA GAG TTT AAT GCT GAA ACA TTC ACC TTC CAT GCA GAT ATA TGC ACA CTT TCT GAG AAG GAG AGA.CAA ATC AAG AAA CAA ACT GCA CTT GTT GAG CTC GTG AAA CAC AAG CCC AAG GCA ACA AAA GAG CAA CTG AAA GCT GTT ATG GAT GAT TTC GCA GCT TTT GTA GAG AAG TGC TGC AAG GCT GAC GAT AAG GAG ACC TGC TTC GCC GAG GAG GGT AAA AAA CTT GTT GCT GCA AGT CAA GCT GCC TTA GGC TTA TAA CATCTACATTTAAAAGCATCTCAGCCTACCATGAGAATA AGAGAAAGAAATGAAGATCAAAAGCTTATTCATCTGTTTTCTTTTTCGTTGGTG TTTTAATCATTTTGCCTCTTTTCTCTGTGCTTCAATTAATAAAAAATGGAAAGAA TCTAA

wherein the 5' to 3' strand, beginning with the amino terminus is shown, and wherein the abbreviations are defined as in claim 4.

- 9. A plasmid having the capability of replication in a prokaryotic or eukaryotic organism, comprising a deoxyribo nucleotide sequence coding for human serum albumin.
- 10. A plasmid as claimed in claim 8 having the capability of replication in a prokaryotic organism, comprising a human serum albumin or human preproserum albumin gene as claimed in any one of claims 1 to 8.

- 11. A plasmid as claimed in claim 9 or claim 10 having the capability of replication in a prokaryotic organism of the genus <u>Escherichia</u>.
- 12. The plasmid of claim 10 designated pGX401 (deposited in <u>E. coli</u> HB101 at the U.S. Dept. of Agriculture Northern Regional Research Center, Peoria, Illinois under accession No. NRRL B-15784) and mutants thereof encoding human serum albumin.

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- 13. A microorganism transformed by a plasmid as10 claimed in any one of claims 9 to 12.
 - 14. A microorganism as claimed in claim 13 of the genus Escherichia.
 - 15. A microorganism as claimed in claim 14 of the species coli.
- 16. A method of producing prepro-human serum albumin which comprises cultivating on an aqueous nutrient medium containing assimilable sources of carbon, nitrogen and essential minerals and growth factors, under prepro-human serum albumin-producing conditions,
- a prokaryotic organism as claimed in claim 13 transformed by a plasmid capable of replicating in said organism and having a deoxyribonucleotide sequence coding for prepro-human serum albumin, and recovering the prepro-human serum albumin so produced.
- 25 17. A method as claimed in claim 16 wherein the prokaryotic organism is E. coli.

- 18. A method as claimed in claim 17 wherein the prokaryotic organism is transformed by a plasmid substantially similar to plasmid pGX401 as claimed in claim 11.
- 5 19. E. coli strain NRRL No. 15784 (pGX401) or a mutant thereof containing a human prepro-human serum albumin gene.

CLAIMS FOR THE DESIGNATED STATE: AT

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- 1. A process for preparing a gene coding for human serum albumin (HSA) which comprises obtaining HSA mRNA from HSA-producing cells, in vitro synthesis of complementary DNA (cDNA) using said mRNA as a template and conversion of said cDNA to the double-stranded form.
- 2. A process as claimed in claim 1 wherein said gene codes for prepro-human serum albumin.
- 3. A process as claimed in claim 1 wherein said gene comprises the following deoxyribonucleotide

 10 sequence which corresponds to the indicated amino acid sequence:

Ala Val His Lys Ser Glu GAY GCX CAY AAM QRS GAM GTX GCX Gly Gl u Phe Lys Asp Leu His Arg CAY LGN TTY AAM GAY YTZ GGX GAM Leu Val Phe Lys Ala Asn GAM AAY TTY AAM GCX YTZ GTX YTZ GlnTyr Leu Ile Ala Phe Ala Gln ATH GCX TTY GCX CAM TAY YTZ CAM Val His Glu Asp Gln Cys Pro Phe CAM TGY CCX TTY GAM GAY CAY GTX Thr Val Val Lys Leu Asn Glu GTX ACX GAM AAM YTZ GTX AAY GAM Ala Val ASD Ala Lys Thr Cys Phe TTY GCX AAM ACX TGY GTX GCX GAY Cys Glu Asn Asp Gl u Ser Ala TGY GAY AAM GAM QRS GCX GAM AAY Phe Gly Asp Leu His Thr Leu TTY GGX GAY QRSYTZ CAY ACX YTZ Thr Val Ala Th r Leu Cys Lys Leu Y T Z AAMYTZ TGY ACX GTX GCX ACX Gl y Glu Met Glu Thr Tyr Arg LGN GAM ACX TAY GGX GAM ATG GCX Lys Gln Glu Ala Asp Cys Cys TGY GCX AAM CAM GAM CCX GAY TGY

Glu Cys Glu Arg Asn Phe Leu GAM LGN AAY GAM TGY TTY YTZ CAM His Lys Asp Asp Asn Pro Asn Leu C A Y A A M G A Y G A Y A A Y C C X A A Y Y T Z Arg Leu Val Arg Pro Glu CCX LGN YTZ GTX LGN CCX GAM GTX Asp Val Met Cys Thr Ala Phe GAY GTX ATG TGY ACX GCX TTY CAY Asp Asn Glu Glu Thr Phe Leu GAY AAY GAM GAM ACX TTY YTZ AAM Lys Tyr Leu Tyr Glu Ile Ala Arg AAM TAY YTZ TAY GAM ATH GCX LGN Phe His Pro Tyr Thr Ala LGN CAY CCX TAY TTY ACX GCX CCX Leu Leu Phe Phe Ala Lys GAM YTZ YTZ TTY TTY GCX AAM LGN Tyr Lys Ala Ala Phe Thr Glu Cys TAY AAM GCX GCX TTY ACX GAM TGY Gln Ala Ala Lys Asp Ala TGY GCX CAM GCX GAY AAM GCX GCX Cys Leu Lys Phe Pro Leu Asp TGY YTZ TTY CCX AAM YTZ GAY GAM Leu Arg Asp Glu Gly Lys Ala Ser Y T Z L G N G A Y G A M G G X A A M G C X Q R S Lys Gln Ala Arg Leu Lys QRS GCX AAM CAM LGN YTZ AAM TGY Ser Gln Leu Lys Phe Gly GCX QRS YTZ CAM AÂM TTY GGX GAM Phe Lys Ala Ala Trp Ala Val L G N G C X T T Y A A M G C X T G G G C X G T X Arq Leu Ser Gln Arg Phe Pro GCX LGN YTZ QRS CAM LGN TTY CCX Glu Phe Ala Ala Glu Val AAM GCX GAM TTY GCX GAM GTX QRS Asp Lys Phe Val Thr Leu AAM TTY GTX ACX GAY YTZ ACX AAM His Thr Glu Cys His Cys GTX CAY ACX GAM TGY TGY CAY GGX

Leu Leu Glu Cys Ala Asp Asp Asp GAY YTZ YTZ GAM TGY GCX GAY GAY Leu Ala Ala Asp Lys Tyr L G N G C X G A Y Y T Z G C X A A M T A Y A T H Cys Glu Asn Gln Asp Ser Ile Ser TGY GAM AAY CAM GAY QRS ATH QRS Ser Lys Leu Lys Glu Cys Cys QRS AAM YTZ AAM GAM TGY TGY GAM Pro Leu Phe Glu Lys Ser AAM CCX YTZ TTY GAM AAM QRS CAY Ala Glu Ile Val Glu Asn TGY ATH GCX GAM GTX GAM AAY GAY Glu Met Pro Ala Asp Phe Pro Ser G A M A T G C C X G C X G A Y T T Y C C X Q R S Ala Phe Ala Val Asp Phe Val Glu TTY GCX GTX GAY TTY GTX GAM QRS Lys Asp Val Cys Lys Asn Tyr AAM GAY GTX TGY AAM AAY TAY GCX Gly Glu Ala Phe Leu Lys Asp Val GAM GCX AAM GAY GTX TTY YTZ GGX Tyr Phe Phe Glu Tyr Ala ATG TTY TTY TAY GAM TAY GCX LGN Val His Pro Tyr Asp Ser Va l L G N C A Y C C X G A Y T A Y Q R S G T X G T X Leu Len Leu Ala Arg Leu Lys YTZ YTZ YTZ LGN YTZ GCX AAM ACX Glu Thr Thr Leu Glu Lvs TAY GAM ACX ACX YTZ GAM AAM TGY Ala Cys Ala Ala Pro His Asp TGY GCX GCX GCX GAY CCX CAY GAM Tyr Ala Lys Val Phe Asp TGY TAY GCX AAM GTX TTY GAY GAM Pro Val Lys Pro Glu Glu Pro TTY AAM CCX CCX GTX GAM GAM CCX

Ile⁴ Gln Asn Phe Ile Lys Gln Asn Cys C A M A A Y T T Y A T H A A M C A M A A Y T G Y Glu Leu Phe Glu Gln Leu Gly Glu G A M Y T Z T T Y G A M C A M Y T Z G G X G A M Lys Phe Gln Asn Ala Leu TAY AAM TTY CAM AAY GCX YTZ TTY Thr Tyr Arq Lys Lys Val GTX LGN TAY ACX AAM AAM GTX CCX Thr Leu Ser Pro Thr Leu CAM YTZ QRS ACX CCX ACX YTZ GTX Arg Gly Val Ser Asn Leu Lvs GAM GTX QRS LGN AAY YTZ GGX AAM Val Gly Ser Lys Cys Cys Lys His G T X G G X Q R S A A M T G Y T G Y A A M C A Y Glu Ala Lys Arg Met Pro CCX GAM GCX AAM LGN ATG CCX TGY Ala Glu Asp Tyr Leu Ser Val Val G C X G A M G A Y T A Y Y T Z Q R S G T X G T X Asp Gln Asn Leu Cys Val Leu Y T Z A A Y C A M Y T Z T G Y G T X Y T Z C A Y Lys Thr Pro Val Ser Asp GAM AAM ACX CCX GTX QRS GAY LGN Glu Thr Thr Lys Cys Cys G T X A C X A A M T G Y T G Y A C X G A M Q R S Gly Leu Val Asn Arg Arg Pro Gly Phe Y T Z G T X A A Y L G N L G N C C X G G X T T Y Ser Ala Leu Glu Val Asp Glu Thr QRS GCX YTZ GAM GTX GAY GAM ACX Val Pro Lys Glu Phe Asn Tyr TAY GTX CCX AAN GAM TTY AAY GCX Phe Ala Thr Phe Thr His Asp GAM ACX TTY ACX TTY CAY GCX GAY Gl u Lys Cys Thr Leu Ser ATH TGY ACX YTZ QRS GAM AAM GAM

Gl n Ile Lys Glu Thr Lys LGN CAM ATH AAM AAM GAM ACX GCX Glu Val Leu Val Lys His Lys YTZ GTX GAM YTZ GTX AAM CAY AAM Glu Leu Pro Lys Ala Thr Lys Glu CCX AAM GCX ACX AAM GAM GAM YTZ Val Asp Phe Lvs Ala Met Asp AAM GCX GTX ATG GAY GAY TTY GCX Phe Val Glu Lys Cys Cys Lys GCX TTY GTX GAM AAM TGY TGY AAM Thr Cys Gl u Phe Ala Asp Asp Lys GCX GAY GAY AAM GAM ACX TGY TTY Lys Glu Glu Gly Lys Leu Val GCX GAM GAM GGX AAM AAM YTZ GTX Gly Val Glu Ala Leu Ala Ser GCX GCX QRS GAM GCX GTX YTZ GGX Leu YTZTAA

wherein, the 5' to 3' strand, beginning with the amino terminus and the amino acids for which each triplet codes are shown, and wherein the abbreviations have the following standard meanings:

. A is deoxyadenyl

T is thymidyl

G is deoxyguanyl

C is deoxycytosyl

X is A, T, C or G

Y is T or C

When Y is C, Z is A, T, C or G

When Y is T, Z is A or G

H is A, T or C

Q is T or A

When Q is T, R is C and S is A, T, C or G

When Q is A, R is G and S is T or C

M is A or G L is A or C When L is A, N is A or G When L is C, N is A, T, C or G GLY is glycine ALA is alanine VAL is valine LEU is leucine ILE is isoleucine SER is serine THR is threonine PHE is phenylalanine TYR is tyrosine TRP is tyryptophan CYS is cysteine MET is methionine ASP is aspartic acid GLU is glutamic acid LYS is lysine ARG is arginine HIS is histidine PRO is proline GLN is glutamine ASN is asparagine

4. A process as claimed in claim 2 wherein said gene comprises the following deoxyribonucleotide sequence:

Trp Met Lys Val Thr Phe ATG AAM TGG GTX ACX TTY Ser Phe Leu Ile Leu Leu ATH QRS YTZ YTZ TTY YTZ Gly Ala Tyr Ser Arg Ser Ser QRS QRS GCX TAY QRS LGN GGX Asp Phe Ala His Val Arg Arg GTX TTY LGN LGN GAY GCX CAY AAM

Phe His Arg Lys Ala Glu Val Ser QRS GAM GTX GCX CAY LGN TTY AAM Asp Leu Gly Glu Glu Asn Phe Lys G A Y Y T Z G G X G A M G A M A A Y T T Y A A M Ala Phe Leu Ile Val Leu GCX YTZ GTX YTZ ATH GCX TTY GCX Phe Cys Pro Leu Gln Gln Tyr CAM TAY YTZ CAM CAM TGY CCX TTY Val Leu Val Lys His Gl u Asp GAM GAY CAY GTX AAM YTZ GTX AAY Phe Ala Lys Thr Glu Val Glu GAM GTX ACX GAM TTY GCX AAM ACX Ala Glu Glu Ser Asp Val Ala TGY GTX GCX GAY GAM QRS GCX GAM Leu Ser Lys Asp Cys. AAY TGY GAY AAM QRS YTZ CAY ACX Leu Phe Gly Asp Lys Leu Cys Thr Y T Z T T Y G G X G A Y A A M Y T Z T G Y A C X Cys Glu Thr Arg Leu Thr Ala GTX GCX ACX YTZ LGN GAM ACX TAY Cys Cys Glu Ala Asp Met GGX GAM ATG GCX GAY TGY TGY GCX Glu Asn Pro Arg Glu Gln AAM CAM GAM CCX GAM LGN AAY GAM Gln His Asp Lys Leu Phe TGY TTY YTZ CAM CAY AAM GAY GAY Asn Pro Asn Leu Pro Arg Leu Val A A Y C C X A A Y Y T Z C C X L G N Y T Z G T X Val Met Val Asp Glu Arg Pro LGN CCX GAM GTX GAY GTX ATG TGY Glu Glu Asn His Asp Ala Phe ACX GCX TTY CAY GAY AAY GAM GAM Lys Tyr Leu Lys Phe Leu ACX TTY YTZ AAM AAM TAY YTZ TAY Glu Ile Ala Arg Arg His Pro Tyr GAM ATH GCX LGN LGN CAY CCX TAY Ala Glu Leu Leu Phe Thr Pro TTY ACX GCX CCX GAM YTZ YTZ TTY Lys Arg Tyr Lys Ala Ala TTY GCX AAM LGN TAY AAM GCX GCX Thr Glu Ala Gln Cys Cys TTY ACX GAM TGY TGY GCX CAM GCX Lys Ala Ala Cys Leu Phe GAY AAM GCX GCX TGY YTZ TTY CCX Glu Leu Asp Leu Asp Arg Lvs AAM YTZ GAY GAM YTZ LGN GAY GAM Gln Lys Ala Ser Ser Ala Lys GGX AAM GCX QRS QRS GCX AAM CAM Leu Lys Cys Ala Ser Arg LGN YTZ AAM TGY GCX QRS YTZ CAM Phe Gly Glu Arg Ala Phe Lys Lys AAM TTY GGX GAM LGN GCX TTY AAM Val Ala Arg Leu Ala Trp Ala GCX TGG GCX GTX GCX LGN YTZ QRS Ala Glu Gln Phe Pro Lys Arg CAM LGN TTY CCX AAM GCX GAM TTY Phe Val Thr Glu Val Ser Lys GCX GAM GTX QRS AAM TTY GTX ACX Val His Thr Thr Lys Leu GAY YTZ ACX AAM GTX CAY ACX GAM Cys Cys His Gly Asp Leu Leu Glu T G Y T G Y C A Y G G X G A Y Y T Z Y T Z G A M Asp Ala Asp Arg Ala Asp TGY GCX GÂY GÂY LGN GCX GÂY YT2 Glu Asn Ile Cys Lys Tyr GCX AAM TAY ATH TGY GAM AAY CAM Lys Leu Lys Ser Ser Ser Ile GAY QRS ATH QRS QRS AAM YTZ AAM

 $|\psi_{ij}\rangle = |\psi_{ij}\rangle \frac{2\pi}{\pi} = |\psi_{ij}\rangle \frac{2\pi}{\pi}$

Cys Cys Glu Lys Pro Leu GAM TĜY TĜY GAM AÃM CCX YTZ TTY Glu Lys Ser His Cys Ile Ala Glu G A M A A M Q R S C A Y T G Y A T H G C X G A M Glu Asn Asp Gl u Met Pro GTX GAM AAY GAY GAM ATG CCX GCX Phe Pro Ser Phe Ala Val GAY TTY CCX QRS TTY GCX GTX GAY Val Glu Ser Lys Asp Val TTY GTX GAM QRS AAM GAY GTX TGY Lys Glu Asn Tyr Ala Ala Asp AAM AAY TAY GCX GAM GCX AAM GAY Gly Met Phe Phe Val Phe Leu GTX TTY YTZ GGX ATG TTY TTY TAY Glu Tyr Ala Arg Arg His Pro Asp G A M T A Y G C X L G N L G N C A Y C C X G A Y Val Ser Val Leu Leu Leu TAY QRS GTX GTX YTZ YTZ YTZ LGN Glu Th r Lys Thr Tyr Y T Z G C X A A M A C X T A Y G A M A C X A C X Cys Ala · Ala Glu Lys Cys YTZ GAM AAM TGY TGY GCX GCX His Ala G1 u Cys Tyr Pro GAY CCX CAY GAM TGY TAY GCX AAM Asp Glu Phe Lys Pro Phe GTX TTY GAY GAM TTY AAM CCX CCX Phe Gln Asn Glu Glu Pro GTX GAM GAM CCX CAM AAY TTY AT 8 Gl u Gln Leu Phe Cys Asn AAM CAM AAY TGY GAM YTZ TTY GAM Gln Leu Gly Glu Tyr Lys Phe Gln CAMYTZGGXGAMTAYAAMTTYCAM Phe Val Arq Tyr Ala Leu AAY GCX YTZ TTY GTX LGN TAY ACX Pro Gln Leu Lys Lys Val Pro Gln Leu Ser Thr AAMAAMGTX CCX CAMYTZ QRS ACX Gl u Val Ser Val Thr Leu CCX ACX YTZ GTX GAM GTX QRS LGN Asn Leu Gly Lys Val Gly Ser Lys A A Y Y T Z G G X A A M G T X G G X Q R S A A M Val Gly Ser Lys Pro Glu Al a His Lys Cys Lys TGY TGY AAM CAY CCX GAM GCX AAM Ala Glu Asp Cys Pro Met LGN ATG CCX TGY GCX GAM GAY TAY Gln Leu Val Val Leu Asn Ser YTZ QRS GTX GTX YTZ AAY CAM YTZ Cys Val Leu His Glu Lys Thr Pro T G Y G T X Y T Z C A Y G A M A A M A C X C C X Val Arg Thr Lys Ser Asp GTX QRS GAY LGN GTX ACX AAM TGY Val Asn Glu Ser Leu Cys Thr TGY ACX GAM ORS YTZ GTX AAY LGN Ser Phe Ala Leu Arg Pro Gly Phe Ser Ala Leu Glu L G N C C X G G X T T Y Q R S G C X Y T Z G A M Tyr Val Pro Asp Glu Thr Val GTX GAY GAM ACX TAY GTX CCX AAM Thr Phe Ala Glu Phe Asn GAM TTY AAY GCX GAM ACX TTY ACX Ala Asp Ile Cys Thr Phe His TTY CAY GCX GAY ATH TGY ACX YTZ Arg Ile Glu Gl n Glu Lys QRS GAM AAM GAM LGN CAM ATH AAM Thr Ala Leu Val Glu Leu Glu Lys AAM GAM ACX GCX YTZ GTX GAM YTZ Lys Ala Lys Pro His Lys GTX AAM CAY AAM CCX AAM GCX ACX Leu Ala Val Met Lys Glu Glu Lys AAM GAM GAM YTZ AAM GCX GTX ATG Ala Phe Val Phe Ala Asp Asp GAY GAY TTY GCX GCX TTY GTX GAM

Lys Cys Cys Ala Asp Lys Asp AĀM TĞY TĞY AÁM GCX GA^TY GAYAAM Thr Cys Phe Gl y Ala Glu Glu G A M A C X T G Y TTYGCX GAM GAM Lys Leu Val Ala Ala Ser AAM YTZ GTX GCX GCX QRS GAM Ala Val Gly Leu Leu G C X G T X Y T Z G G X Y T Z

wherein the 5' and 3' strand, beginning with the amino terminus, and the amino acids for which each triplet codes are shown, and wherein the abbreviations are defined as in claim 3.

5. A process as claimed in claim 3 wherein said gene comprises the following deoxyribonucleotide sequence:

GAT GCA CAC AAG AGT GAG GTT GCT CAT CGG TTT AAA GAT TTG GGA GAA GAA AAT TTC AAA GCC TTG GTG TTG ATT GCC TTT GCT CAG TAT CTT CAG CAG TGT CCA TTT GAA GAT CAT GTA AAA TTA GTG AAT GAA GTA ACT GAA TTT GCA AAA ACA TGT GTT GCT GAT GAG TCA GCT GAA AAT TGT GAC AAA TCA CTT CAT ACC CTT TTT GGA GAC AAA TTA TGC ACA GTT GCA ACT CTT CGT GAA ACC TAT GGT GAA ATG GCT GAC TGC TGT GCA AAA CAA GAA CCT GAG AGA AAT GAA TGC TTC TTG CAA CAC AAA GAT GAC AAC CCA AAC CTC CCC CGA TTG GTG AGA CCA GAG GTT GAT GTG ATG TGC ACT GCT TTT CAT GAC AAT GAA GAG ACA TTT TTG AAA AAA TAC TTA TAT GAA ATT GCC AGA AGA CAT CCT TAC TTT TAT GCC CCG GAA CTC CTT TTC TTT GCT AAA AGG TAT AAA GCT GCT TTT ACA GAA TGT TGC CAA GCT GCT GAT AAA GCT GCC TGC CTG TTG CCA AAG CTC GAT GAA CTT CGG GAT GAA GGG AAG GCT TCG TCT GCC AAA CAG AGA CTC AAG TGT GCC AGT CTC CAA AAA TTT GGA GAA AGA GCT TTC AAA GCA TGG GCG GTG GCT CGC CTG AGC CAG AGA TTT CCC AAA GCT GAG TTT GCA GAA GTT TCC AAG TTA GTG ACA GAT CTT ACC AAA GTC CAC ACG GAA TGC TGC CAT GGA GAT CTG CTT GAA TGT GCT GAT GAC AGG GCG GAC CTT GCC AAG TAT ATC TGT GAA AAT CAA GAT TCG ATC TCC AGT AAA CTG AAG GAA TGC TGT GAA

AAA CCT CTG TTG GAA AAA TCC CAC TGC ATT GCC GAA GTG GAA AAT GAT GAG ATG CCT GCT GAC TTG CCT TCA TTA GCT GCT GAT TTT GTT GAA AGT AAG GAT GTT TGC AAA AAC TAT GCT GAG GCA AAG GAT GTC TTC CTG GGC ATG TTT TTG TAT GAA TAT GCA AGA AGG CAT CCT GAT TAC TCT GTC GTG CTG CTG CTG AGA CTT GCC AAG ACA TAT GAA ACC ACT CTA GAG AAG TGC TGT GCC GCT GCA GAT CCT CAT GAA TGC TAT GCC AAA GTG TTC GAT GAA TTT AAA CCT CCT GTG GAA GAG CCT CAG AAT TTA ATC AAA CAA AAT TGT GAG CTT TTT GAG CAG CTT GGA GAG TAC AAA TTC CAG AAT GCG CTA TTA GTT CGT TAC ACC AAG AAA GTA CCC CAA GTG TCA ACT CCA ACT CTT GTA GAG GTC TCA AGA AAC CTA GGA AAA GTG GGC AGC AAA TGT TGT AAA CAT CCT GAA GCA AAA AGA ATG CCC TGT GCA GAA GAC TAT CTA TCC GTG GTC CTG AAC CAG TTA TGT GTG TTG CAT GAG AAA ACG CCA GTA AGT GAC AGA GTC ACC AAA TGC TGC ACA GAA TCC TTG GTG AAC AGG CGA CCA TGC TTT TCA GCT CTG GAA GTC GAT GAA ACA TAC GTT CCC AAA GAG TTT AAT GCT GAA ACA TTC ACC TTC CAT GCA GAT ATA TGC ACA CTT TCT GAG AAG GAG AGA CAA ATC AAG AAA CAA ACT GCA CTT GTT GAG CTC GTG AAA CAC AAG CCC AAG GCA ACA AAA GAG CAA CTG AAA GCT GTT ATG GAT GAT TTC GCA GCT TTT GTA GAG AAG TGC TGC AAG GCT GAC GAT AAG GAG ACC TGC TTT GCC GAG GAG GGT AAA AAA CTT GTT GCT GCA AGT CAA GCT GCC TTA GGC TTA TAA wherein the 5' to 3' strand, beginning with the amino terminus is shown, and wherein the abbreviations are defined as in claim 3.

6. A process as claimed in claim 4 wherein said gene comprises the following deoxyribonucleotide .sequence:

ATG AAG TGG GTA ACC TTT ATT TCC CTT CTT TTT CTC TTT AGC
TCG GCT TAT TCC AGG GGT GTG TTT CGT CGA GAT GCA CAC AAG
AGT GAG GTT GCT CAT CGG TTT AAA GAT TTG GGA GAA GAA AAT
TTC AAA GCC TTG GTG TTG ATT GCC TTT GCT CAG TAT CTT CAG
CAG TGT CCA TTT GAA GAT CAT GTA AAA TTA GTG AAT GAA GTA
ACT GAA TTT GCA AAA ACA TGT GTT GCT GAT GAG TCA GCT GAA

AAT TGT GAC AAA TCA CTT CAT ACC CTT TTT GGA GAC AAA TTA TGC ACA GTT GCA ACT CTT CGT GAA ACC TAT GGT GAA ATG GCT GAC TGC TGT GCA AAA CAA GAA CCT GAG AGA AAT GAA TGC TTC TTG CAA CAC AAA GAT GAC AAC CCA AAC CTC CCC CGA TTG GTG AGA CCA GAG GTT GAT GTG ATG TGC ACT GCT TTT CAT GAC AAT GAA GAG ACA TIT TIG AAA AAA TAC TIA TAT GAA ATI GCC AGA AGA CAT CCT TAC TTT TAT GCC CCG GAA CTC CTT TTC TTT GCT AAA AGG TAT AAA GCT GCT TTT ACA GAA TGT TGC CAA GCT GCT GAT AAA GCT GCC TGC CTG TTG CCA AAG CTC GAT GAA CTT CGG GAT GAA GGG AAG GCT TCG TCT GCC AAA CAG AGA CTC AAG TGT GCC AGT CTC CAA AAA TTT GGA GAA AGA GCT TTC AAA GCA TGG GCG GTG GCT CGC CTG AGC CAG AGA TTT CCC AAA GCT GAG TTT GCA GAA GTT TCC AAG TTA GTG ACA GAT CTT ACC AAA GTC CAC ACG GAA TGC TGC CAT GGA GAT CTG CTT GAA TGT GCT GAT GAC AGG GCG GAC CTT GCC AAG TAT ATC TGT GAA AAT CAA GAT TCG ATC TCC AGT AAA CTG AAG GAA TGC TGT GAA AAA CCT CTG TTG GAA AAA TCC CAC TGC ATT GCC GAA GTG GAA AAT GAT GAG ATG CCT GCT GAC TTG CCT TCA TTA GCT GCT GAT TTT GTT GAA AGT AAG GAT GTT TGC AAA AAC TAT GCT GAG GCA AAG GAT GTC TTC CTG GGC ATG TTT TTG TAT GAA TAT GCA AGA AGG CAT CCT GAT TAC TCT GTC GTG CTG CTG AGA CTT GCC AAG ACA TAT GAA ACC ACT CTA GAG AAG TGC TGT GCC GCT GCA GAT CCT CAT GAA TGC TAT GCC AAA GTG TTC GAT GAA TTT AAA CCT CCT GTG GAA GAG CCT CAG AAT TTA ATC AAA CAA AAT TGT GAG CTT TTT GAG CAG CTT GGA GAG TAC AAF TTC CAG AAT GCG CTA TTA GTT CGT TAC ACC AAG AAA GTA COO CAA GTG TCA ACT CCA ACT CTT GTA GAG GTC TCA AGA AAC CTA GGA AAA GTG GGC AGC AAA TGT TGT AAA CAT CCT GAA GCA AAA AGA ATG CCC TGT GCA GAA GAC TAT CTA TCC GTG GTC CTG AAC CAG TTA TGT GTG TTG CAT GAG AAA ACG CCA GTA AGT GAC AGA GTC ACC AAA TGC TGC ACA GAA TCC TTG GTG AAC AGG CGA CCA TGC TTT TCA GCT CTG GAA GTC GAT GAA ACA TAC GTT CCC AAA GAG TTT AAT GCT GAA ACA TTC ACC TTC CAT GCA GAT ATA TGC ACA CTT TCT GAG AAG GAG AGA CAA ATC AAG AAA CAA ACT GCA CTT GTT GAG CTC GTG AAA CAC AAG CCC AAG GCA ACA AAA GAG CAA CTG AAA GCT GTT ATG GAT GAT TTC GCA GCT TTT GTA GAG AAG TGC TGC AAG GCT GAC GAT AAG GAG ACC TGC TTT GCC GAG GAG GGT AAA AAA CTT GTT GCT GCA AGT CAA GCT GCC TTA GGC TTA TAA wherein the 5' to 3' strand, beginning with the amino terminus is shown, and wherein the abbreviations are defined as in claim 3.

7. A process as claimed in claim 6 wherein said gene is comprised in the following deoxyribonucleotide sequence:

5 '

TCTCTTCTGTCAACCCCACGCCTTTGGCACA ATG AAG TGG GTA ACC TTT ATT TCC CTT CTT TTT CTC TTT AGC TCG GCT TAT TCC AGG GGT GTG TTT CGT CGA GAT GCA CAC AAG AGT GAG GTT GCT CAT CGG TTT AAA GAT TTG GGA GAA GAA AAT TTC AAA GCC TTG GTG TTG ATT GCC TTT GCT CAG TAT CTT CAG CAG TGT CCA TTT GAA GAT CAT GTA AAA TTA GTC AAT GAA GTA ACT GAA TTT GCA AAA ACA TGT GTT GCT GAT GAG TCA GCT GAA AAT TGT GAC AAA TCA CTT CAT ACC CTT TTT GGA GAC AAA TTA TGC ACA GTT GCA ACT CTT CGT GAA ACC TAT GGT GAA ATG GCT GAC TGC TGT GCA AAA CAA GAA CCT GAG AGA AAT GAA TGC TTC TTG CAA CAC AAA GAT GAC AAC CCA AAC CTC CCC CGA TTG GTG AGA CCA GAG GTT GAT GTG ATG TGC ACT GCT TTT CAT GAC AAT GAA GAG ACA TTT TTG AAA AAA TAC TTA TAT GAA-ATT GCC AGA AGA CAT CCT TAC TTT TAT GCC CCG GAA CTC CTT TTC TTT GCT AAA AGG TAT AAA GCT GCT TTT ACA GAA TGT TGC CAA GCT GCT GAT AAA GCT GCC TGC CTG TTG CCA AAG CTC GAT GAA CTT CGG GAT GAA GGG AAG GCT TCG TCT GCC AAA CAG AGA CTC AAG TGT GCC AGT CTC CAA AAA TTT GGA GAA AGA GCT TTC AAA GCA TGG GCG GTG GCT CGC CTG AGC CAG AGA TTT CCC AAA GCT GAG TTT GCA GAA GTT TCC AAG TTA GTG ACA GAT CTT ACC AAA GTC CAC ACG GAA TGC TGC CAT GGA GAT CTG CTT GAA TGT GCT GAT GAC AGG GCG GAC CTT GCC AAG TAT ATC TGT GAA AAT CAA GAT TCG ATC TCC AGT AAA CTG AAG GAA TGC TGT GAA AAA CCT CTG TTG GAA AAA TCC CAC TGC ATT GCC GAA GTG GAA AAT GAT GAG ATG CCT GCT GAC TTG CCT TCA TTA GCT GCT GAT TTT GTT GAA AGT AAG GAT GTT TGC AAA AAC TAT GCT GAG GCA AAG GAT GTC TTC CTG GGC ATG TTT TTG TAT GAA TAT GCA AGA AGG CAT CCT GAT TAC TCT GTC GTG CTG CTG CTG AGA CTT GCC AAG ACA TAT GAA ACC ACT CTA GAG AAG TGC TGT GCC GCT GCA GAT CCT CAT GAA TGC TAT GCC AAA GTG TTC GAT GAA TTT AAA CCT CCT GTG GAA GAG CCT CAG AAT TTA ATC AAA CAA AAT TGT GAG CTT TTT GAG CAG CTT GGA GAG TAC AAA TTC CAG AAT GCG CTA TTA GTT CGT TAC ACC AAG AAA GTA CCC CAA GTG TCA ACT CCA ACT CTT GTA GAG GTC TCA AGA AAC CTA GGA AAA GTG GGC AGC AAA TGT TGT AAA CAT CCT GAA GCA AAA AGA ATG CCC TGT GCA GAA GAC TAT CTA TCC GTG GTC CTG AAC CAG TTA TGT GTG TTG CAT GAG AAA ACG CCA GTA AGT GAC AGA GTC ACC AAA TGC TGC ACA GAA TCC TTG GTG AAC AGG CGA CCA TGC TTT TCA GCT CTG GAA GTC GAT GAA ACA TAC GTT CCC AAA GAG TTT AAT GCT GAA ACA TTC ACC TTC CAT GCA GAT ATA TGC ACA CTT TCT GAG AAG GAG AGA CAA ATC AAG AAA CAA ACT GCA CTT GTT GAG CTC GTG AAA CAC AAG CCC AAG GCA ACA AAA GAG CAA CTG AAA GCT GTT ATG GAT GAT TTC GCA GCT TTT GTA GAG AAG TGC TGC AAG GCT GAC GAT AAG GAG ACC TGC TTC GCC GAG GAG GGT AAA AAA CTT GTT GCT GCA AGT CAA GCT GCC TTA GGC TTA TAA CATCTACATTTAAAAGCATCTCAGCCTACCATGAGAATA **AGAGAAAGAAATGAAGATCAAAAGCTTATTCATCTGTTTTCTTTTTCGTTGGTG** TTTTAATCATTTTGCCTCTTTTCTCTGTGCTTCAATTAATAAAAAATGGAAAGAA

wherein the 5' to 3' strand, beginning with the amino terminus is shown, and wherein the abbreviations are defined as in claim 3.

- 8. A process for preparing a plasmid encoding human serum albumin which comprises inserting a deoxy-ribonucleotide sequence coding for human serum albumin into a plasmid having the capability of replication in a prokaryotic or eukaryotic organism.
- 9. A process as claimed in claim 8 wherein the deoxyribonucleotide sequence coding for human serum albumin is prepared by a process as claimed in any one of claims 1 to 7.

10. A process as claimed in claim 8 or claim 9 wherein the deoxyribonucleotide sequence coding for human serum alubmin is inserted into a plasmid having the capability of replication in a prokaryotic organism of the genus Escherichia.

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- 11. The process of claim 10 wherein the deoxyribo-nucleotide sequence of claim 8 is inserted at the Pst I site of plasmid pBR322 so as to prepare plasmid pGX401.
- 12. A process for preparing a microorganism containing a gene coding for human serum albumin which comprises transforming a microorganism with a plasmid capable of replicating in said microorganism and including said gene.
- 13. A process as claimed in claim 12 wherein said plasmid is prepared by a process as claimed in any one of claims 8 to 11.
- 14. A microorganism transformed by a plasmid containing a deoxyribonucleotide sequence as defined in any one of claims 3 to 7.
 - 15. A microorganism as claimed in claim 14 of the genus Escherichia.
- 16. A method of producing prepro-human serum albumin which comprises cultivating on an aqueous nutrient medium containing assimilable sources of carbon, nitrogen and essential minerals and growth factors, under prepro-human serum albumin-producing conditions, a prokaryotic organism transformed by a plasmid capable of replicating in said organism and having a deoxyribonucleotide sequence coding for

prepro-human serum albumin, and recovering the preprohuman serum albumin so produced.

- 17. A method as claimed in claim 16 wherein the prokaryotic organism is \underline{E} . \underline{coli} .
- 5 18. A method as claim in claim 17 wherein the prokaryotic organism is transformed by a plasmid substantially similar to plasmid pGX401.

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19. <u>E. coli</u> strain NRRL No. 15784 (pGX401), or a mutant thereof containing a human prepro-human serum albumin gene.

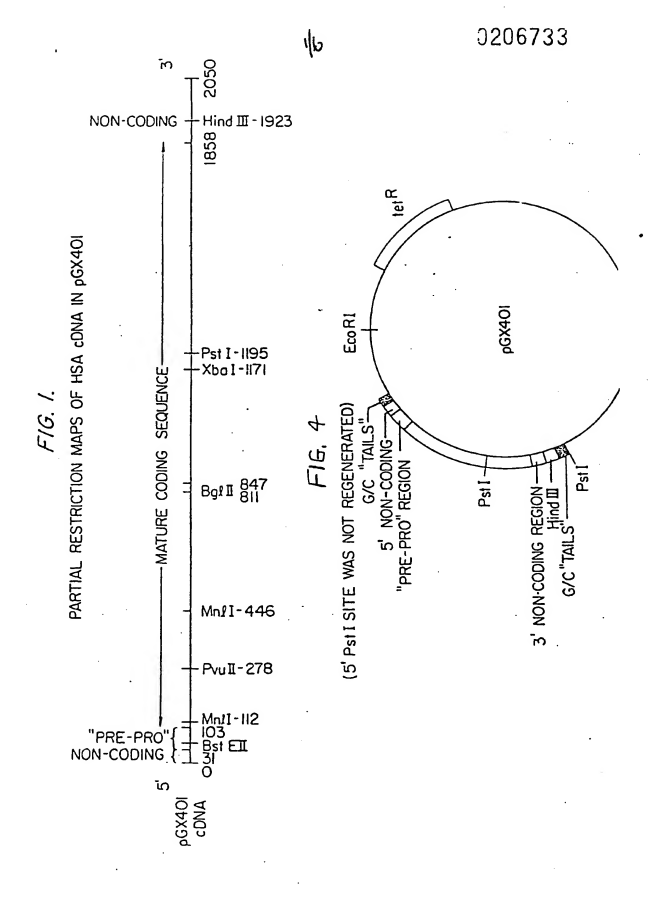


Figure 2

Complete Nucleotide Sequence of the HSA Insert In Clone pGX401

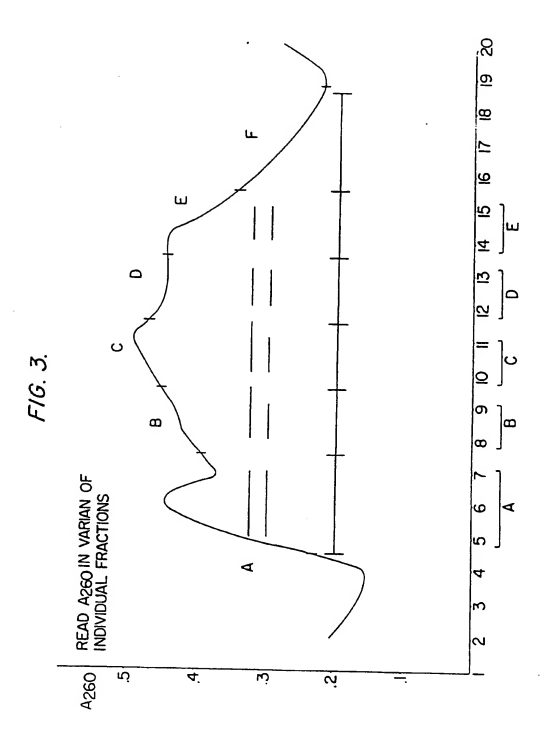
5' TO	CTCTI	CTGI	CAAC	CCCA	cecc	TTTC	GCAC	Йe		s Tr	p Va G GT		
						- Dre	428						-> l
Thr	Phe TTT	Ile ATT	Ser TCC	Leu	Leu	Phe	Leu	Phe	Ser	Ser	Ala GCT	Tyr	Ser
Arg	Glv	Val	Phe	Arg	Arg	Asp	Ala	His	Lys	Ser	Glu GAG	Val	Ala
His	Arg	Phe	Lys	Asp	Leu	Gly	Glu	Glu	Asn	Phe	Lys AAA	Ala	Leu
Val	Leu	Ile	Ala	Phe	Ala	Gln	Tyr	Leu	Gln	Gln	Cys TGT	Pro	Phe
Glu	Asp	His	Val	Lys	Leu	Val	Asn	Glu	Val	Thr	Glu	Phe	Ala
GAA	GAT	CAT	GTA	AAA	TTA	GTC	AAT	GAA	GTA	ACT	GAA	TTT	GCA
Lys	Thr	Cys	Val	Ala	Asp	Glu	Ser	Ala	Glu	Asn	Cys	Asp	Lys
AAA	ACA	TGT	GTT	GCT	GAT	GAG	TCA	GCT	GAA	TAA	TGT	GAC	AAA
Ser	Leu	His	Thr	Leu	Phe	Gly	Asp	Lys	Leu	Cys	Thr	Val	Ala
TCA	CTT	·CAT	ACC	CTT	TTT	GGA	GAC	AAA	TTA	TGC	ACA	GTT	GCA
Thr	Leu	Arg	Glu	Thr	Tyr	Gly	Glu	Met	Ala	Asp	Cys	Cys	Ala
ACT	CTT	CGT	GAA		TAT	GGT	GAA	ATG	GCT	GAC	TGC	TGT	GCA
Lys	Gln	Glu	Pro	Glu	Arg	ASN	Glu	Cys	Phe	Leu	Gln	His	Lys
AAA	CAA	GAA	CCT	GAG	AGA	AAT	GAA	TGC	TTC	TTG	CAA	CAC	AAA
Asp	Asp	Asn	Pro	Asn	Leu	Pro	Arg	Leu	Val	Arg	Pro	Glu	Val
GAT	GAC	AAC	CCA	AAC	CTC	CCC	CGA	TTG	GTG	AGA		GAG	GTT
Asp	Val	Met	Cys	Thr	Ala	Phe	His	Asp	Asn	Glu	Glu	Thr	Phe
GAT	GTG	ATG	TGC	ACT	GCT	TTT	CAT	GAC	AAT	GAA	GAG		TTT
Leu	Lys	Lys	Tyr	Leu	Tyr	Glu	Ile	Ala	Arg	Arg	His	Pro	Tyr
TTG	AAA	AAA	TAC	TTA	TAT	GAA	ATT	GCC	AGA	AGA	CAT	CCT	TAC
Phe TTT	Thr TAT	Ala GCC	Pro CCG	Glu GAA	Leu CTC	Leu CTT	Phe TTC	Phe TTT	Ala	Lys	Arg AGG	Tyr TAT	Lys AAA

Figure 2 (continued)

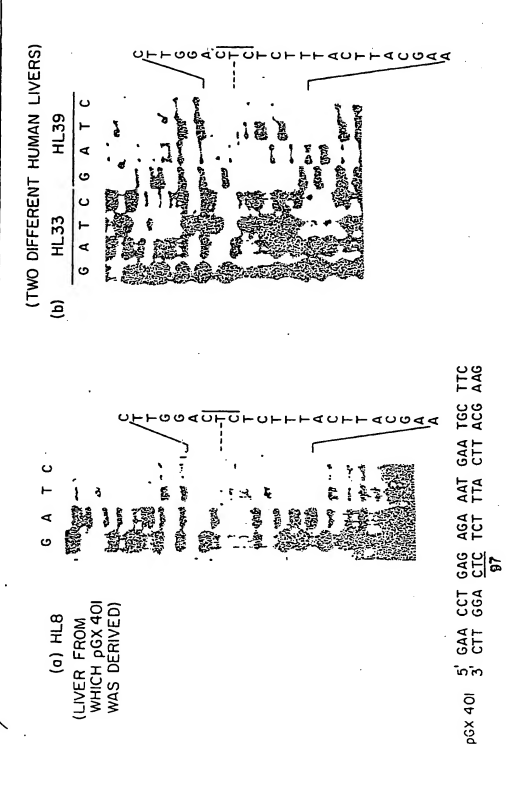
Ala Ala Phe Thr Glu Cys Cys Ala Gln Ala Asp Lys Ala Ala GCT GCT TTT ACA GAA TGT TGC CAA GCT GCT GAT AAA GCT GCC Cys Leu Phe Pro Lys Leu Asp Glu Leu Arg Asp Glu Gly Lys TGC CTG TTG CCA AAG CTC GAT GAA CTT CGG GAT GAA GGG AAG Ala Ser Ser Ala Lys Gln Arg Leu Lys Cys Ala Ser Leu Gln GCT TCG TCT GCC AAA CAG AGA CTC AAG TGT GCC AGT CTC CAA Lys Phe Gly Glu Arg Ala Phe Lys Ala Trp Ala Val Ala Arg AAA TTT GGA GAA AGA GCT TTC AAA GCA TGG GCG GTG GCT CGC Leu Ser Gln Arg Phe Pro Lys Ala Glu Phe Ala Glu Val Ser CTG AGC CAG AGA TTT CCC AAA GCT GAG TTT GCA GAA GTT TCC Lys Phe Val Thr Asp Leu Thr Lys Val His Thr Glu Cys Cys AAG TTA GTG ACA GAT CTT ACC AAA GTC CAC ACG GAA TGC TGC His Gly Asp Leu Leu Glu Cys Ala Asp Asp Arg Ala Asp Leu CAT GGA GAT CTG CTT GAA TGT GCT GAT GAC AGG GCG GAC CTT Ala Lys Tyr Ile Cys Glu Asn Gln Asp Ser Ile Ser Ser Lys GCC AAG TAT ATC TGT GAA AAT CAA GAT TCG ATC TCC AGT AAA Leu Lys Glu Cys Cys Glu Lys Pro Leu Phe Glu Lys Ser His CTG AAG GAA TGC TGT GAA AAA CCT CTG TTG GAA AAA TCC CAC Cys Ile Ala Glu Val Glu Asn Asp Glu Met Pro Ala Asp Phe TGC ATT GCC GAA GTG GAA AAT GAT GAG ATG CCT GCT GAC TTG Pro Ser Phe Ala Val Asp Phe Val Glu Ser Lys Asp Val Cys CCT TCA TTA GCT GCT GAT TTT GTT GAA AGT AAG GAT GTT TGC Lys Asn Tyr Ala Glu Ala Lys Asp Val Phe Leu Gly Met Phe AAA AAC TAT GCT GAG GCA AAG GAT GTC TTC CTG GGC ATG TTT Phe Tyr Glu Tyr Ala Arg Arg His Pro Asp Tyr Ser Val Val TTG TAT GAA TAT GCA AGA AGG CAT CCT GAT TAC TCT GTC GTG Leu Leu Leu Arg Leu Ala Lys Thr Tyr Glu Thr Thr Leu Glu CTG CTG CTG AGA CTT GCC AAG ACA TAT GAA ACC ACT CTA GAG Lys Cys Cys Ala Ala Ala Asp Pro His Glu Cys Tyr Ala Lys AAG TGC TGT GCC GCT GCA GAT CCT CAT GAA TGC TAT GCC AAA Val Phe Asp Glu Phe Lys Pro Pro Val Glu Glu Pro Gln Asn GTG TTC GAT GAA TTT AAA CCT CCT GTG GAA GAG CCT CAG AAT Phe Ile Lys Gln Asn Cys Glu Leu Phe Glu Gln Leu Gly Glu TTA ATC AAA CAA AAT TGT GAG CTT TTT GAG CAG CTT GGA GAG

Figure 2 (continued)

Tyr Lys Phe Gln Asn Ala Leu Phe Val Arg Tyr Thr Lys Lys TAC AAA TTC CAG AAT GCG CTA TTA GTT CGT TAC ACC AAG AAA Val Pro Gln Leu Ser Thr Pro Thr Leu Val Glu Val Ser Arg GTA CCC CAA GTG TCA ACT CCA ACT CTT GTA GAG GTC TCA AGA Asn Leu Gly Lys Val Gly Ser Lys Cys Cys Lys His Pro Glu AAC CTA GGA AAA GTG GGC AGC AAA TGT TGT AAA CAT CCT GAA Ala Lys Arg Met Pro Cys Ala Glu Asp Tyr Leu Ser Val Val GCA AAA AGA ATG CCC TGT GCA GAA GAC TAT CTA TCC GTG GTC Leu Asn Gln Leu Cys Val Leu His Glu Lys Thr Pro Val Ser CTG AAC CAG TTA TGT GTG TTG CAT GAG AAA ACG CCA GTA AGT Asp Arg Val Thr Lys Cys Cys Thr Glu Ser Leu Val Asn Arg GAC AGA GTC ACC AAA TGC TGC ACA GAA TCC TTG GTG AAC AGG Arg Pro Gly Phe Ser Ala Leu Glu Val Asp Glu Thr Tyr Val CGA CCA TGC TTT TCA GCT CTG GAA GTC GAT GAA ACA TAC GTT Pro Lys Glu Phe Asn Ala Glu Thr Phe Thr Phe His Ala Asp CCC AAA GAG TTT AAT GCT GAA ACA TTC ACC TTC CAT GCA GAT Ile Cys Thr Leu Ser Glu Lys Glu Arg Gln Ile Lys Lys Glu ATA TGC ACA CTT TCT GAG AAG GAG AGA CAA ATC AAG AAA CAA Thr Ala Leu Val Glu Leu Val Lys Bis Lys Pro Lys Ala Thr ACT GCA CTT GTT GAG CTC GTG AAA CAC AAG CCC AAG GCA ACA Lys Glu Glu Leu Lys Ala Val Met Asp Asp Phe Ala Ala Phe AAA GAG CAA CTG AAA GCT GTT ATG GAT GAT TTC GCA GCT TTT Val Glu Lys Cys Cys Lys Ala Asp Asp Lys Glu Thr Cys Phe GTA GAG AAG TGC TGC AAG GCT GAC GAT AAG GAG ACC TGC TTC Ala Glu Glu Gly Lys Lys Leu Val Ala Ala Ser Glu Ala Val GCC GÁG GAG GGT AÃA AÃA CTT GTT GCT GCA AGT CAA GCT GCC Leu Gly Leu STOP TTA GGC TTA TAA CATCTACATTTAAAAGCATCTCAGCCTACCATGAGAATA AGAGAAAGAAATGAAGATCAAAAGCTTATTCATCTGTTTTCTTTTTCGTTGGTG TTTTAATCATTTTGCCTCTTTTCTCTGTGCTTCAATTAATAAAAAATGGAAAGAA TCTAA



F/G. \mathcal{S} . DIRECT NUCLEOTIDE SEQUENCE DETERMINATION FROM mRNA





EUROPEAN SEARCH REPORT

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